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http://dx.doi.org/10.24382/3801 University of Plymouth

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TITANIUM DIOXIDE NANOPARTICLE UPTAKE ACROSS THE ISOLATED PERFUSED INTESTINE OF RAINBOW TROUT: PHYSIOLOGICAL MECHANISMS AND A COMPARISON WITH CACO-2 CELLS

By

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A thesis submitted to the Plymouth University in

partial fulfilment for the degree of

DOCTOR OF PHILOSOPHY

School of Biomedical and Biological Sciences

Faculty of Science and Technology

July 2013

Titanium dioxide nanoparticle uptake across the isolated perfused intestine of rainbow trout: physiological mechanisms and a comparison with Caco-2 cells

Aliaa R. Al-Jubory

Abstract

The wide use of nanoscale materials in food and health care products raises the concern of their possible uptake across the gastrointestinal tract, but very limited data are available on their uptake kinetics, and the potential hazards for humans. In this study, the uptake mechanism of titanium dioxide (TiO_2) across the isolated perfused fish intestine and human intestinal Caco-2 cells were evaluated. The in vitro preparation of the whole gut sac and the isolated perfused intestine of rainbow trout were performed using both bulk and nano TiO₂ in a concentration of 1 mg l^{-1} for up to 4 h. The results showed that the Ti from both bulk and TiO₂ NPs were mainly accumulated in the mid and hind intestine, with 80% or more of the accumulation in the mucosa rather than the underlying muscularis. Perfused intestines showed a saturable, time-dependent accumulation of the Ti from TiO₂ and the uptake of Ti from exposure to NPs was faster than that of the bulk form. The uptake of Ti from exposure to TiO₂ NPs increases 10 fold when the CO_2 in the gas mixture was lowered to 0.5%. Subsequently, further investigation on the mechanisms of uptake of TiO₂ was applied using different kinds of inhibitors. Adding 10 mmol l⁻¹ cyanide did not stop Ti uptake from TiO₂ exposures, and 100 μ mol l⁻¹ vanadate (ATPase inhibitor) caused a 2.8 fold reduction in the net uptake rate of Ti for the TiO₂ NP exposure. Luminal additions of 120 IU ml⁻¹ nystatin (endocytosis inhibitor) blocked the uptake of Ti from both bulk and TiO₂ NPs treatments. The results indicate that Ti accumulation from TiO₂ exposures was sensitive

to both nystatin and vanadate; the former suggesting that there is an endocytosis involvement in the uptake of TiO₂ across the intestinal epithelium. Human intestinal Caco-2 cell showed a steady, saturable and time-dependent accumulation of Ti over 24 h exposures to 1 mg 1^{-1} TiO₂ (for all forms of TiO₂). A scanning electron microscope study indicated the appearance of the particles underneath the cells, increasing the evidence of the Ti uptake from different forms of TiO₂ by Caco-2 cells. Both nystatin and vanadate increase the accumulation of TiO₂ which suggests interference of these drugs with endocytic pathways. All the data in the thesis demonstrates Ti uptake across the intestinal epithelium from TiO₂ exposures involving CO₂-dependent and nystatinsensitive mechanisms. The results in this thesis have contributed to some understanding on the behaviour, uptake and effects of the TiO₂ NPs across the intestine; and highlight the possible dietary hazard of the NPs to human health.

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Abbreviations

Abbreviation	Glossary	
%	Percent	
°C	Degree Centigrade	
μg	Micrograms	
μl	Microliter	
μm	Micrometer	
μmol	Micromole	
A431	Human epidermal cell line	
A549	Human lung adenocarcinoma epithelial cells	
Ag	Silver	
ANOVA	Analysis of variance	
ARPE-19	Human retinal pigment epithelial cells	
ATP	Adenosine triphoshate	
BAL	Bronchoalveolar lavage	
BCA	Bicinchoninic acid	
BEAS-2B	Human bronchial epithelial cells	
BSI	Backscatter image	
C ₆₀	Buckminsterfullerene	
Ca	Calcium	
CaCl ₂ ·2H ₂ O	Calcium chloride dihydrate	
Caco-2	Human colon adenocarcinoma cell line	
CARS	Coherent anti-Stokes Raman scattering	
СВ	Carbon black	
Cd	Cadmium	
CH ₃ Hg	Methylmercury	
Cl	Chloride	
cm	Centimetre	
cm ²	Square centimetre	

CNT	Carbon nanotubes	
CO_2	Carbon dioxide	
CO ₃	Carbon trioxide	
Cu	Copper	
DEPs	Diesel exhaust nanoparticles	
DIDS	4,4'-diisothiocyano stilbene-2,2'-disulfonic acid	
DLVO	Derjaguin, Landau, Verwey and Overbeek	
DMEM	Dulbecco's modified eagle medium	
DNA	Deoxyribonucleic acid	
D.O.M	Dissolved Organic Matter	
DPBS	Dulbecco's phosphate buffered saline	
E171	Titanium dioxide	
EDTA	Ethylene diamine tetraacetic acid	
EDS	Energy dispersive spectrophotometry	
EGTA	Ethylene glycol tetraacetic acid	
FBS	Fetal bovine serum	
Fe ³⁺	Ferric iron	
Fig	Figure	
g	Gram	
GIT	Gastrointestinal tract	
GSH	Glutathione	
GTP	Guanosine triphosphate	
H^+	Hydrogen ion	
h	Hour	
HaCaT	Human keratinocyte cell line	
HBDH	Alpha-hydroxybutyrate dehydrogenase	
HCl	Hydrochloric acid	
HCO ₃	Bicarbonate	
H & E	Haematoxylin and eosin stain	
HEPES	4-(2-hydroxyethyl)-1-piperazineethanesulfonic acid	

Hg	Inorganic mercury	
His	Histidine	
ICP-OES	Inductivity coupled plasma-optical emission spectrome	
IU	International unit	
Κ	Potassium	
KCl	Potassium chloride	
KCN	Potassium cyanide	
Kg	Kilogram	
kGy	Kilogray	
kHz	Kilohertz	
1	Litre	
LC ₅₀	Median lethal concentration	
LD ₅₀	Median lethal dose	
LDH	Lactate dehydrogenase	
М	Molar	
M cells	Microfold cells	
MD	Minimal Davis medium	
MDCK	Madin darby canine kidney cells	
Mg	Magnesium	
mg	Milligram	
MgCO ₃	Magnesium carbonate	
MgSO ₄	Magnesium sulphate	
Milli-Q	Ultra-pure ion free water	
min	Minute	
ml	Millilitre	
mM	Millimolar	
mm	Millimetre	
mmol	Millimole	
Mn	Manganese	
MnO ₂	Manganese dioxide	

mol	Mole	
mosm	Milliosmole	
mW	milliwatt	
MWCNTS	Multi-walled carbon nanotubes	
n	Number of observations	
Na	Sodium	
NaCl	Sodium chloride	
NADH	Nicotine amide adenine dinucleotide	
NaHCO ₃	Sodium bicarbonate	
NaH ₂ PO ₄ [·] 2H ₂ O	Sodium dihydrogen phosphate dehydrate	
Na ₃ Vo ₄	Sodium orthovanadate	
ng	Nanogram	
nm	Nanometer	
NMs	Nanomaterials	
nmol	Nanomole	
NPs	Nanoparticles	
NRR	Neutral red retention	
NTA	Nanoparticle tracking analysis	
O_2	Oxygen	
OsO4	Osmium tetroxide	
Р	Statistical probability	
P25	Titanium dioxide nanoparticles	
PC-3M	Human prostate carcinoma cells	
pCO_2	Partial pressure of carbon dioxide	
PM	Particulate matter	
рН	Potential hydrogen ion	
PHFK	Primary human foreskin keratinocytes	
PLGA	Poly (lactic-co-glycolic acid)	
PTRF	Polymerase 1 and transcript release factor	
QDs	Quantum dots	

r	Round	
RBC	Red blood cell	
REACH	Registration, Evaluation, Authorisation of Chemicals	
ROS	Reactive oxygen species	
sCT	Salmon calcitonin	
sec	Second	
SEI	Secondary electron image	
S.E.M	Standard error of mean	
SEM	Scanning electron microscopy	
SiO ₂	Silicon dioxide	
SSA	Specific surface area	
SWCNTS	Single-walled carbon nanotubes	
TBARS	Thiobarbituric acid reactive substance	
TEM	Transmission electron microscopy	
Ti	Titanium	
TiO ₂	Titanium dioxide	
Ti(OH) ₄	Titanium hydroxide	
TiO(OH) ₂	Metatitanic acid	
USL	Unstirred layer	
UV	Ultraviolet	
UVA	Ultraviolet A	
vit	Vitamins	
WBC	White blood cell	
Zn	Zinc	
ZnO	Zinc oxide	

Publications (Please refer to Appendix I):

- Handy RD, Al-Bairuty G, Al-Jubory A, Ramsden CS, Boyle D, Shaw BS and Henry TB 2011. Effects of manufactured nanomaterials on fishes: a target organ and body systems physiology approach. *Journal of Fish Biology*, 79, 821-853.
- **Al-Jubory AR** and Handy RD 2012. Uptake of titanium from TiO₂ nanoparticles exposure in the isolated perfused intestine of rainbow trout: nystatin, vanadate, and novel CO₂-sensitive components. *Nanotoxicology*, DOI: 10.3109/17435390. 2012.735268.

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- **Al-Jubory AR**, Handy RD and Henry T 2009. The physiological effects of nanoparticles. University of Plymouth, 3rd April 2009, Plymouth, UK.
- Al-Jubory AR, Handy RD and Henry T 2012. Mechanisms uptake of TiO₂ nanoparticles across the isolated perfused intestine of rainbow trout (*Oncorhynchus mykiss*). UoP Nano Group Meeting, University of Plymouth, 3rd Annual Meeting, 10th May 2012, Plymouth, UK.
- Al-Jubory AR, Handy RD and Henry T 2012. Uptake of TiO₂ nanoparticles across the isolated perfused intestine of rainbow trout: nystatin, vanadate, and novel CO₂-sensitive componenets. Ecotoxicology Research and Innovation Centre (ERIC), University of Plymouth, 2nd Annual Conference, 13th July 2012, Plymouth, UK.

Poster presentations and conferences abstracts (Please refer to Appendix II):

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- **Al-Jubory A** and Handy RD 2010. Uptake of TiO₂ nanoparticles across the isolated perfused intestine of rainbow trout (*Oncorhynchus mykiss*). Nanotoxicology 2010, Edinburgh, UK, 2nd-4th June 2010, abstract book, pp. 70, No. p 95.
- Al-Jubory A and Handy RD 2010. Uptake of TiO₂ nanoparticles across the isolated perfused intestine of rainbow trout (*Oncorhynchus mykiss*). Society of Environmental Toxicology and Chemistry UK branch annual meeting, London, 13-14th September 2010. Abstract book, SETAC.29, p 25.
- **Al-Jubory A** and Handy RD 2011. Uptake of TiO₂ nanoparticles across the isolated perfused intestine of rainbow trout (*Oncorhynchus mykiss*). 3rd NanoImpactNet Conference, "Building a Bridge from NanoImpactNet to Nanomedical Research", 14-17th February 2011, Lausanne, Switzerland, abstract book, 4.2.6, p 70.

- **Al-Jubory AR** and Handy RD, 2011. Mechanisms uptake of TiO₂ nanoparticles across the isolated perfused intestine of rainbow trout (*Oncorhynchus mykiss*). Society of Environmental Toxicology and Chemistry UK Branch Annual Meeting, "Polluted Planet: Sustaining Ecosystems and Biodiversity", London, 12-13th September 2011. Abstract book, SETAC.113.
- **Al-Jubory AR** and Handy RD, 2012. Uptake of TiO₂ nanoparticles across the isolated perfused intestine of rainbow trout: nystatin, vanadate, and novel CO₂-sensitive componenets. The Postgraduate Society Annual Conference, Plymouth University, 26th June 2012, Plymouth, UK.
- Gitrowski C, **Al-Jubory AR** and Handy RD 2012. Uptake and retention of TiO₂ nanoparticles in Caco-2 cells implications for human health. Centre for Research in Translational Biomedicine (CRTB) Research Day, University of Plymouth, 2nd Annual Conference, Plymouth, 4th July 2012.
- Gitrowski C, **Al-Jubory AR**, Handy RD 2013. The gastrointestinal uptake of TiO₂ nanoparticles and associated toxicity. Society of Environmental Toxicology and Chemistry, SETAC Europe 23rd annual meeting, "Building a better future: Responsible innovation and environmental protection", 12-16th May 2013, Glasgow, UK. Abstract book, No. TU058, p229-230.

Acknowledgments

I would like to express my sincere gratitude to my first supervisor Professor Richard Handy for his help, advices, encouragement and immense knowledge along the way from the beginning up to the completion of this study. I would like to extend my sincere thanks to my second supervisor Dr. Ted Henry for his support and encouragement throughout my study.

I would like also to thanks the technical colleagues, especially William Vevers, Andy Atfield and Benjamin Eynon for their technical support where needed. Special thanks go out to Dr. Andrew Fisher for his help with ICP-OES analysis and a big thanks to Peter Bond for his help in TEM and SEM work. Christopher Ramsden is thanked for metal analysis advices and answering my questions, Genan Al-Bairuty for histology advices and help. I would like to acknowledge Constantinos Gitrowski who worked with me as a team. We worked together in the Caco-2 cell experiment. I would like also to express my sincere appreciation to all my colleagues and friends in the School of Biomedical and Biological Sciences who answered my request for help, especially Helena Reinardy, Sahar Karieb and Sahar Al Kutby. Professor Waleed Al-Murrani is thanked for his support and encouragement. I would like to express my gratitude to all those people, though their names are not listed, who supported me in one way or another.

Many thanks to the Ministry of Higher Education and Scientific Research-Republic of Iraq for providing the financial support and Iraqi cultural attaché in London for their kind support. Finally, I would like to express my heartfelt thanks to all my family especially my beloved parents for their blessing and supports, sisters and brothers for their encouragement. Lastly, my special heartfelt thanks go to my husband Mudhafar who supported me along the time of my study, my children Noor and Mohammed for their patient and encouragement.

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Author's declaration

At no time during the registration for the degree of Doctor of Philosophy has the author been registered for any other University award without prior agreement of the Graduate Committee. This study was fully funded by the Ministry of Higher Education and Scientific Research of Republic of Iraq.

Word count of main body of thesis: 49,286 (56,947 including references)

Signed -----

Date-----

Dedication

This work is dedicated to my beloved husband, Mudhafar Al-Zandee and my two children, Noor and Mohammed



Chapter 1

General Introduction

1.1 Introduction to nanoparticles and nanotechnology

Nanotechnology and the use of nano-scale materials is a relatively new area of science and technology which has become a significant worldwide industry, and is growing rapidly (www.nano.org.uk). Nanoparticles (NPs) are defined as particles with at least one dimension less than 100 nm (Masciangioli and Zhang, 2003; Roco, 2003) and the range size of these NPs should be between a lower limit of 1 nm and upper limit of 100 nm (Lövestam et al., 2010). This definition is not absolute, according to toxicological studies which have included work on aggregates of particles with dimensions of a few hundred nanometers (Handy and Shaw, 2007; Handy et al., 2008a). Clearly for NPs, the primary size should be considered (e.g., the diameter of a single particle). In addition, the sizes of aggregates of NPs, which can be several hundred nanometers or more, and the distribution of particle sizes present in the material also need to be considered. In mammalian toxicology particles sizes (PM, particulate matter) have been defined as coarse particles (diameter between 10 μ m and 2.5 μ m, PM 10-2.5), fine particles (2.5 μ m or less, PM 2.5), or ultrafine particles (< 0.1 μ m, PM 0.1), so NPs could be regarded as ultrafine particles or smaller (Handy et al., 2008a).

There are a wide variety of nanomaterials (NMs) and particles at the nanoscale range, because these NMs are purposefully designed for a particular application it may be difficult to classify these materials into chemical groups, and the use of the product (risk according to how products are used) may be more important for the exposure aspects of risk assessment of these materials (Hansen et al., 2008). Considerable effort has therefore focused on the classification of NMs according to the industrial aspect they are used in, for example: domestic appliances, electronics and computers, food and food packaging, health and fitness which includes (personal care products, cosmetics, sunscreen and sporting goods), home and garden, water treatment technology and environmental remediation (Masciangioli and Zhang, 2003; Roco, 2003; Aitken et al., 2006; Hansen et al., 2008).

There is also another classification of NMs proposed by Hansen et al. (2007) which broadly classifies NMs, not in terms of products, but according to how the material is incorporated into the product, including three main categories: whether NMs are the bulk of the material in the product, whether they exist on the surface (surface coated products), or forming part of the structure (NPs matrix within the product). These main categories could be also divided into subcategories; the bulk of the material may consist of just one type of NMs or more than two different types (Hansen et al., 2007). For the surface coated products, the surface coated with NMs consist of the same structure material like the bulk, surface covers with un-patterned film of NMs thickness different from the substrate material, or the surface is consist of pattern film on a substrate (the nanoscale material is either the film or the pattern). The NPs in the third category may be present in four subcategories depending on the environment; the NPs may be bound to another solid structure surface, suspended in liquid, dispersed into a solid, or the NPs can be suspended in air (airborne NPs) (Hansen et al., 2007).

Risk assessment of new materials is traditionally based on a chemicals approach, and so there has also been debate about how to chemically classify NPs and NMs. The many different chemical structures may justify the classification of these materials into some broad chemical categories initially (Jortner and Rao, 2002; Rao, 2004; Handy et al., 2008a; Stone et al., 2010). These chemical classifications of the NPs are shown in Table 1.1. Table 1.1 Chemical categories of nanoparticles.

Chemical categories	Sub-category	Materials
Carbon based materials	Nanotuhas	Single-walled carbon nanotubes "SWCNTS"
	Nallotubes	Multi-walled carbon nanotubes "MWCNTS"
	Fullerenes	C _{60,} buckyballs
Metal based materials	Metals	Gold, silver, iron
	Metal oxides	Titanium dioxide, zinc oxide, silicon oxide
	Metal sulphides	Zinc sulphides, copper sulphides
	Nitrides of metals	Silicon, cadmium, tellurium
Organic	Polymers	Polyethyleneglycol, latex
	Dendrimers	Polyamidoamide
Composites	polymer nanocomposites	Multicomponent nanomaterial
	quantum dots	Multicomponent nanomaterial

1.2 Physico-chemical properties of nanoparticles

One of the main concerns about nanomaterials is that their novel properties will lead to new or unknown toxic effects. It is therefore critical to consider the physico-chemistry of NPs in toxicology (Lead and Wilkinson, 2006; Handy et al., 2008a; Landsiedel et al., 2010). There is much research to do on this aspect, but factors of concern include surface area, shape, crystal structure, surface reactivity and how aggregation chemistry influences toxicity (Handy et al., 2008a, Stone et al., 2010, Von der Kammer et al., 2012). NPs can show aggregation behaviour which partly depends on the large surface area proportional to their volume. Particles may form agglomerates (loosely bound particles) which bonded together by weak forces, such as Vander Waals forces, electrostatic forces and surface tension. Alternatively, particles may also form aggregates (groups of strongly connected particles) that are not easily redispersed mechanically (Handy et al., 2008a). These forces are explained by the DLVO theory which is named after Derjaguin and Landau, (1941), and Verwey and Overbeek, (1948). When the NPs aggregate in a solution they should be follow this theory i.e., remain as single particles in a solution, or form aggregates via particle-particle, particle-cluster and cluster-cluster interactions (Handy et al., 2008a). The theory also predicts that ionic strength (e.g., changes in salinity), and the presence of cations such as Ca^{2+} (e.g., changes in water hardness), or other charged particles will influence this aggregation behaviour (Borm et al., 2006; Handy et al., 2008a, Von der Kammer et al., 2012). There are many physico-chemical properties to investigate in toxicology studies. So far, particular emphasis has been given to the surface area issue. The behaviour of NPs can be influenced by the increased specific surface area (SSA) proportional to the small size of NPs, and this may increase the bioavailability and toxicity of the NPs (Handy et al., 2008a). According to Oberdörster et al. (2007) the dose effect of NMs can be better explained based on the SSA rather than concentration, suggesting the former may be more important in the toxicity of NPs. Similar arguments have been made about surface reactivity, since it is this surface that presents toxic chemicals to the organism. A huge increase in a surface: volume ratio, and hence the presence of more surface molecules should result in more intrinsic toxicity of the materials (Donaldson and Tran, 2004), and for some particles there is evidence that toxicity may be a function of particle size rather than mass concentration (Oberdörster, 2000). Surface charge also is an issue, a study by Schaeublin et al. (2011) showed that gold NPs sized at 1.5 nm, either positively charged, neutral and negatively charged, cause disrupted to the morphology of human keratinocyte cell line (HaCaT); with cell death through apoptosis induced by charged NPs and cell necrosis by neutral NPs. Other studies on invertebrates also emphasized the differences in acute toxicity of NP is dependent on sample preparation and therefore the presence of small or larger aggregates in the test medium (Lovern and Klaper, 2006). The toxicity of NPs may be further influenced by the shape of the materials. Hamilton et al. (2009) showed that anatase nanobelt increased markers of the inflammation detected in bronchoalveolar lavage (BAL) of mice in comparison to those detected with TiO₂ nanospheres in treated animals.

1.3 Routes of exposure

The exposures of humans to manufactured NPs in the environment provide different routes to penetrate the human body, i.e. the skin (dermal penetration), respiratory tract (inhalation), and gastrointestinal tract (ingestion) (Hoet et al., 2004; Oberdörster et al., 2005b; Yah et al., 2012). Humans could also be exposed to NPs through other routes (medical devices, cosmetics, or clinical procedures using manufactured NMs, Nel et al., 2006; Arora et al., 2012). Another possible route of exposure is by parental administration of NPs in the case of biomedical applications like intravenous, intradermal and peritoneal exposures (Yah et al., 2012).

1.3.1 Skin

The skin is the largest organ of the body and interfaces with the environment. The skin is considered as the largest primary defence of the body against external interferences. The skin is directly exposed to NPs through the application of sunscreen creams, cosmetics, skin care products and drug treatments (Oberdörster et al., 2005b; Hagens et al., 2007). Then the NPs could penetrate into the deeper parts of the skin; and have been localized within the epidermal and dermal layers (e.g., Fullerene-based peptides, Rouse et al., 2006; TiO₂ NPs, Gontier et al., 2008).

1.3.2 Respiratory system

Inhalation is considered as the main route of exposure to NPs in humans and recently researchers found different pathogenic effects after inhalation of different manufactured NPs (Oberdörster et al., 2005a; Donaldson et al., 2006; Lam et al., 2006; Song et al., 2009). An *in vitro* study on the exposure of human bronchial epithelial cells (BEAS-2B) to different concentrations of SWCNT (0.06 mg ml⁻¹, 0.12 mg ml⁻¹ or 0.24 mg ml⁻¹) for 18 h at 37 °C showed that SWCNT caused structural and morphological changes, cellular toxicity, apoptosis and induced an oxidative stress effect in the lung (Shvedova

et al., 2003). Other studies showed that the absorption of NPs by the lung epithelium, primarily through the alveolar region, could reach different internal body systems or organs like the spleen, liver, bone marrow, lymph nodes, heart and brain via blood and lymph (Nemmar et al., 2001; Kreyling et al., 2002; Oberdörster et al., 2005a; Hagens et al., 2007).

1.3.3 Gastrointestinal tract (GIT)

Another important route of exposure to NPs is via ingestion. Gastrointestinal tract exposure to NPs could be happening either via an indirect route through the respiratory system after excretion of inhaled particles by the mucociliary escalator and then ingested into GIT (Oberdörster et al., 2005b; Hagens et al., 2007), or directly ingested via food, water, drugs, cosmetics and drug delivery devices (Oberdörster et al., 2005b; Arora et al., 2012; Yah et al., 2012). There is a wide concern about NPs toxicity through food (Tiede et al., 2008; Fröhlich and Roblegg, 2012). Weir et al. (2012) shows that body exposure to TiO_2 via food was on average between 1-3 mg TiO_2/Kg body weight /day depending on the age and dietary habits considered 36% of these particles in nanoscale range.

Absorption of NPs through the gut epithelium have been shown by some studies and with sometimes with evidence of translocatone into other internal organs (Jani et al., 1990; Borm and Kreyling, 2004; Oberdörster et al., 2005a; Oberdörster et al., 2005b; Chung et al., 2010). Koeneman et al. (2010) showed that TiO_2 NP at 10 µg ml⁻¹ and above could cross the human intestinal cell line (Caco-2) and penetrate through the cells via a transcellular vesicular trafficking pathway without disturbing junctional complexes.

Oral and dietary exposures to TiO_2 NPs *in vivo* show some toxic effects to the GIT in different organisms. A study by Wang et al. (2007) indicates some pathology in liver and kidney (hepatic injury and nephrotoxicity) of mice after single oral gavage of

 TiO_2 NPs (25 or 80 nm) and observed Ti accumulation in spleen, kidney and lung. However, the dietary exposure of TiO_2 NPs showed less toxicity to the GIT and detected Ti accumulation in different organs (e.g., rat, Jani et al., 1994; fish, Ramsden et al., 2009).

1.4 Toxic effects of NPs on different organisms

Studies on the toxic effects of NPs have been applied to different groups of organisms including human, mammals, birds, fish, invertebrates, plants, as well as bacteria. A number of vertebrates (amphibians and reptiles) still lack information on NPs toxicity to date. Some key findings on different groups of organisms are summarised below.

1.4.1 Mammals and birds

There are many studies on NPs toxicity on mammals, and most of these studies have focused on the respiratory toxicity. Pathologies are observed in rodents after intratracheal instillation or aerosol inhalation of different NPs, including the formation of epithelial granuloma, fibrosis and inflammation according to the type of particle used (Bermudez et al., 2004; Lam et al., 2004; Warheit et al., 2004; Warheit et al., 2006). Lam et al. (2004) showed that exposure of mice to 0.5 mg/mouse of SWCNT induced 56% mortality with lung lesions, necrosis, macrophages and granulomas, interstitial and peribronchial inflammation.

Also, there is NP toxicity detected in liver, spleen, and kidney of mice after dietary exposures (Chen et al., 2006; Wang et al., 2006). Mice exposed to copper NPs via oral gavage demonstrated acute toxicity with pathological damage to kidney, liver and spleen compare to bulk copper; and observed colour changes (staining intensity) in both spleen and kidney with the atrophy of spleen after ingesting nano-copper particles (Chen et al., 2006). While the exposure of adult mice to nano-zinc suspensions showed the symptoms of lethargy, nausea, vomiting and diarrhoea (Wang et al., 2006). Liver and heart damage, disturbances of energy and amino acid metabolism was observed after oral administration of TiO_2 NP in rats (Bu et al., 2010).

There is a very limited data on NP exposures to birds. All the studies have been focused on silver NPs with no significant effects shown for different parameters (white blood cells, blood enzymes, embryonic growth and development) in broiler chicken or chicken embryo (Grodzik and Sawosz, 2006; Sawosz et al., 2007; Ahmadi and Kurdestany, 2010; Pineda et al., 2012).

1.4.2 Fish, aquatic and terrestrial invertebrates

For the fishes, most of the studies have been on fresh water fish. There is evidence of a range of different toxic effects with some data on the accumulation of NPs in different organs (Oberdörster, 2004; Oberdörster et al., 2006; Federici et al., 2007; Smith et al., 2007; Ramsden et al., 2009; Wu et al., 2010; Fraser et al., 2011; Shaw and Handy, 2011; Jahanbakhshi et al., 2012). Aqueous exposure of rainbow trout (*Oncorhynchus mykiss*) to different sizes of Ag NPs for 10 days resulted in accumulation of silver in the gills and liver with oxidative stress effects found in the gills (Scown et al., 2010). The accumulation of silver in the liver was twice that found in the gills, which suggests that the important route of exposure and uptake of Ag NPs in fish may be via the gut epithelium (Scown et al., 2010).

There are lots of data available on fresh water invertebrates specially in *Daphnia magna* (Lovern and Klaper, 2006; Oberdörster et al., 2006; Lovern et al., 2007; Roberts et al., 2007; Gao et al., 2009). The exposure of *Daphnia magna* to 0.2-10 mg1⁻¹ of TiO₂ NPs (10-20 nm) for 48h showed 100 % mortality (Lovern and Klaper, 2006). *Daphnia magna* showed acute toxicity and biomodification when exposed to 20 mg1⁻¹ SWCNT (Lipid coated) (1.2 nm) and 100% mortality observed after 48 h (Roberts et al., 2007).

Only a few data have been collected on terrestrial invertebrates (Jemec et al., 2008; Drobne et al., 2009; Wang et al., 2009; Hu et al., 2010; Li et al., 2011). A study
by Li et al. (2011) showed that the exposure of earthworm *Eisenia fetida* to 1000 mg ZnO NP Kg⁻¹ agar for short time (96 h) resulted in 100% mortality. However, using the filter paper method demonstrated highest mortality rate with lower concentrations (50 mg I^{-1}). This mortality seemed to decline with increasing exposure concentration (Li et al., 2011).

One study on the marine invertebrate (*Arenicola marina*) lugworm showed a significant decrease in casting rate, as well as increases in cellular and DNA damage after exposure to 1 g Kg⁻¹ TiO₂ NP through natural sediments for 10 days. However, no significant effects were seen after exposed to 0.03 g Kg⁻¹ SWNT (Galloway et al., 2010).

1.4.3 Plants

Only a few data are available on interaction of NPs with plants (Corredor et al., 2009; Wang et al., 2011; Lee et al., 2012). In a hydroponics study, a slight reduction on root elongation was found in presence of uncoated alumina NPs but not with phenanthrene coated NPs (Yang and Watts, 2005). One study on green algae showed some toxic effects, observed by a growth reduction after the exposure to TiO_2 NPs (Hund-Rinke and Simon, 2006). TiO₂ also induced a positive effect on the growth of spinach, when either administered to the seeds or sprayed onto the leaves (Zheng et al., 2005; Yang et al., 2006).

1.4.4 Bacteria

Toxicological studies of NPs are limited on bacteria, protozoa, and other microbes (Lyon et al., 2005; Adams et al., 2006; Zhu et al., 2006; Zhang et al., 2007; Pelletier et al., 2010). Zhu et al. (2006) showed that the uptake of CNTs by unicellular protozoans induced a dose-dependent growth inhibition. Inorganic NPs (TiO₂, SiO₂ and ZnO) had toxic effects on bacteria (*Escherichia coli* and *Bacillus Subtilis*) represented by

enhancing generation of reactive oxygen species associated with the inhibition of bacterial growth (Adams et al., 2006). In addition, the presence of light was a significant factor for increasing this toxicity (Adams et al., 2006).

1.5 Systemic toxicity

The toxicity of certain NPs could be reaching beyond the site of uptake to the major internal body systems. In most studies, the lethal dose (LD₅₀) may not be estimated because of the difficulty to achieve the high mg levels needed to achieve acute toxicity for many NPs (Handy et al., 2008a). For example, Chen et al. (2006) showed that the LD₅₀ determined for 23.5 nm copper NPs and cupric ions in mice exposed via oral gavage were 413 and 110 mg kg⁻¹ body weight, respectively. Lovern and Klaper (2006) also suggest that the medial lethal concentration (LC₅₀) level for 30 nm TiO₂ is 5.5 mg l⁻¹ in *Daphnia magna*. It is therefore likely that the main concern for NPs will be sublethal effects on the physiology and biochemistry of the animals. Most of the literature on the body systems toxicity of NPs comes from mammalian studies mainly on respiratory exposure, or from *in vitro* cell studies (Handy and Shaw, 2007).

1.5.1 Respiratory system

There is an historic literature on respiratory exposure to ultrafine particles, and arguably, the respiratory route has been investigated the most. Exposure of the respiratory system to NPs has demonstrated lung injury, inflammation and possibly tumour formation in the lung in several studies (Bermudez et al., 2004; Lam et al., 2004; Warheit et al., 2004, 2006). Epithelial cells of the mammalian lung represent a typical mucous epithelial tissue, which is not fundamentally different to the structure of the epithelia of gill or gut of the aquatic organisms, or the body surface of earth worms (Handy et al., 2008b). There is also evidence of the similar epithelial injury observed in fish gill (Federici et al., 2007; Smith et al., 2007; Sun et al., 2009).

1.5.2 Gastrointestinal tract (GIT)

The gut is also an important epithelial tissue, and a potential route of uptake of toxic chemicals. The data on dietary exposure to NPs is currently limited (Panessa-Warren et al., 2006; Arora et al., 2012). However, exposure of mice to 5 g kg⁻¹ of zinc NPs (58 nm) caused severe symptoms of lethargy, vomiting and diarrhoea for the first few days and two mortalities after the first week; the dead mice showed aggregation of Zn particles in the intestine (Wang et al., 2006). Federici et al. (2007) reported that the exposure of rainbow trout to 0.1-1 mg1⁻¹ of TiO₂ NPs (24 nm) for up to 14 days via the water (stress-induced by drinking contaminated water) caused injuries to the intestine including erosion of the villi, as well as fusion and vacuolation of the mucosa. Ramsden et al. (2009) showed an accumulation of Ti in gut after dietary exposure of rainbow trout to 10 or 100 mg kg⁻¹ TiO₂ NPs. The accumulation of arsenic in the stomach and intestine of carp had been significantly increased by 132% in presence of 10 mg 1⁻¹ TiO₂ NPs after 25 days aqueous exposure (Sun et al., 2007). Effects on NPs on nutritional performance are not currently clear, and data on aspects such as growth rate, functions of digestive enzymes and absorption remain to be determined.

1.5.3 Integumentary system (skin)

There are limited *in vivo* studies that have been conducted on the cutaneous toxicity of NPS. Studies on skin cultures, subcutaneous implantation (1 and 4 weeks) in rats exposed to carbon nanofibers and carbon nanotubes (0.1 mg l^{-1}) observed foreign body granuloma (Sato et al., 2005; Yokoyama et al., 2005).

1.5.4 Cardiovascular and immunology system

Once the NP has crossed the lung (gill), gut or skin it will have access to the cardiovascular system and body fluids. There is little data on the cardiotoxicity of NPs. The heart is very sensitive to oxidative stress and low oxygen conditions (e.g., carbon

monoxide exposure, Patel et al., 2004), and this is a concern for NPs that may be oxidising agents. The administration by oral gavages of 5 mg kg⁻¹ TiO₂ NPs (25 and 80 nm) in adult mice produced a significant enhancement of serum lactate dehydrogenase (LDH), and alpha-hydroxybutyrate dehydrogenase (HBDH) suggesting myocardial damage (Wang et al., 2007). The exposure of human red blood cells (cell culture) to 5 μ g ml⁻¹ ultrafine TiO₂ (30-40 nm) for 4-24 h showed by electronic microscope intracellular NPs aggregates (RBCs uptake) with no morphological changes (no cell damage) (Rothen-Rutishauser et al., 2006).

The intraperitoneal injection of male rats with 1 ml of different concentrations (25, 50 and 100 mg l^{-1}) of cerium oxide NPs showed an immune response to the NPs exposure indicated by the increase in number of white blood cells. Subsequently, increasing the exposure with 200 mg l^{-1} of cerium oxide NPs caused severely reduced numbers of white blood cells compared to the control and other lower exposure concentrations (Hamrahi-michak et al., 2012).

1.5.5 Brain and central nervous system

The effects of NPs on the brain and central nervous system are also a concern. The inhalation of 30 nm MnO_2 NP in rats was associated with elevated levels of Mn in the brain, including the frontal cortex, striatum and cerebellum (Elder et al., 2006). An *in vitro* study by Hussain et al. (2006) indicated the ability of manganese NPs to induce adverse effects in neuronal cells, including loss of cell viability, induction of oxidative stress, and dopamine depletion. Studies on trout brain showed pathological deformations including possible aneurisms or swellings on the ventral surface of the cerebellum after exposure to 0.25 mg Γ^1 SWCNT (Smith et al., 2007). Oberdörster (2004) showed a significant elevation of the lipid peroxidation in the brain after exposure of the juvenile largemouth bass to 0.5 mg Γ^1 C₆₀ fullerenes for up to 48 h, but did not include a vehicle control in their experiment.

1.5.6 Internal organs (kidney, liver and spleen) and reproductive system

There is some information on the effects of NPs on internal organs. Exposure of mice to copper NPs (23.5 nm diameter), dosed with different concentrations (108, 158, 232, 341, 501, 736 and 1080 mg kg⁻¹) via oral gavage, induced toxicological effects indicated by significant injuries in the kidney, liver and spleen (Chen et al., 2006). In the kidney, the pathology included swelling of renal glomeruli, degeneration and irreversibly massive necrobiosis of epithelial cells of renal proximal convoluted tubules. In the liver, steatosis around the venae centrals of hepatic tissue was reported and atrophic changes in the spleen (Chen et al., 2006).

In fish, the exposure of rainbow trout to TiO_2 NPs or SWCNT showed minor fatty change and lipidosis in liver cells, and some hepatocytes showed condensed nuclear bodies (apoptotic bodies) (Federici et al., 2007; Smith et al., 2007). After aqueous exposure to 100 and 200 mg l⁻¹ TiO₂ NPs for 20 days in carp, different pathologies was observed in the liver including necrotic and apoptosis hepatocytes with depletion of antioxidant enzymes activities and elevation of liver lipid peroxidation (Hao et al., 2009). Ti accumulations were observed in the liver and spleen after dietary exposure of 10 or 100 mg kg⁻¹ TiO₂ NPs for up to 8 weeks (Ramsden et al., 2009).

Data on the effects of NPs on reproductive behaviours, gamete production, and reproductive success appear to be lacking. *In vitro* study by Komatsu et al. (2008) found effects on Leydig cells, including changes in cell viability and gene expression after expose of mouse Leydig TM3 cells to diesel exhaust NPs (DEPs), TiO₂ and carbon black (CB) NPs.

1.6 Titanium dioxide NPs (TiO₂)

 TiO_2 is a non-combustible and odourless white powder which naturally exists in three mineral structures that include anatase, rutile and brookite crystal forms (Chen and Mao,

2007; Macwan et al., 2011). The anatase and rutile forms are manufactured commercially and both of them have a tetragonal crystal system, whereas the rutile form has a denser arrangement of atoms (Iavicoli et al., 2011). TiO₂ NPs are being used as a white pigment for a wide range of paints, paper, plastic, ceramics, and with the ability to absorb and reflect UV light, it has been used in sunscreens and cosmetics (Aitken et al., 2006; Hansen et al., 2008). These wide applications cause wariness about the ability of TiO₂ NPs to penetrate the human, or animal bodies, by different routes of exposure (Sayes et al., 2006; Wang et al., 2007; Warheit et al., 2007b; Xie et al., 2011) which raise the concern about the toxicity of TiO₂ NPs.

1.6.1 Toxic effects of TiO₂ NPs

Several studies in rodents and aquatic organisms have demonstrated some toxic effects of $TiO_2 NPs$. Examples of *in vivo* and *in vitro* toxicity data are shown in Table 1.2 and 1.3. Ultrafine-TiO₂ particles have been considered as more toxic than fine particles (Oberdörster et al., 1992; Bermudez et al., 2002; Bermudez et al., 2004; Warheit et al., 2007a). There may also be different biological effects of these materials such as inflammation and fibrosis of the lung (Oberdörster et al., 1992; Bermudez et al., 2004; Warheit et al., 2006), or even lung tumours after chronic exposures (Heinrich et al., 1995), see Table 1.2 and 1.3.

Both the anatase and rutile crystal forms of TiO_2 are toxic (Warheit et al., 2007b), but the anatase crystal form is considered as a superior catalysts, a better generator of reactive oxygen species (ROS), and therefore more cytotoxic compared to the rutile particle form (Uchino et al., 2002; Sayes et al., 2006). The data so far suggests that TiO_2 NPs may be toxic via several mechanisms.

Cytotoxicity of different concentrations of TiO_2 NP have been observed after exposure of the human bronchial epithelial cell line (BEAS-2B), which led to cell death, increases of reactive oxygen species (ROS), and decreased glutathione (GSH) (Park et al., 2008). Alterations in thiobarbituric acid substances (TBARS), which indicates general changes in oxidative stress, were observed in tissues from trout after exposure to the $TiO_2 NP$ (Federici et al., 2007). $TiO_2 NPs$ in the anatase form produced the most ROS generation and the largest cytotoxic responses following *in vitro* exposures to human dermal fibroblasts or to human lung epithelial cells (Sayes et al., 2006).

TiO₂ NPs may also be neurotoxic. The exposure of mouse brain microglia cells (BV2 cells) to different concentrations of TiO₂ NPs (P25) stimulated the production of ROS by the microglia through an oxidative burst suggesting interference with mitochondrial energy production (Long et al., 2006). Exposure of mice to different concentrations of TiO₂ NP for 60 days via intragastric administration caused brain damage and a decline in behaviours associated with memory of spatial recognition (Hu et al., 2010).

DNA damage has also been observed in several studies (Gurr et al., 2005; Trouiller et al., 2009; Shukla et al., 2011), see Table 1.2 and 1.3. Vevers and Jha (2008) showed the induction of DNA strand breaks (including oxidative damage to the DNA) after the interactive toxic potential of TiO_2 NPs with ultraviolet radiation (UVA) and also lysosomal membrane integrity using neutral red retention (NRR) assay in an established fish cell line.

The exposure to TiO_2 NPs can influence the uptake of other pollutants. For example, there was a 146% increase in the accumulation of cadmium (Cd) in the presence of TiO_2 NPs compared with a control of Cd exposure alone in carp (Zhang et al., 2007) as well as *Daphnia magna* and *Lumbriculus variegatus* (Hartmann et al., 2012). Some of these effects may relate to the surface properties of NPs. Table 1.2 Toxic effects of TiO₂ NPs in vivo on mammals and aquatic organisms.

TiO ₂ NPs Information	Dose / Time	Exposure Route	Species	Toxic Effects	Author
Ultrafine TiO ₂ average size 21 nm	0.5, 2.0 or 10mg m^{-3} for 6h/day for 5 days a week over 13 weeks	Aerosol inhalation	Mice, rats and hamsters	Pulmonary inflammation in rats and mice (increased number of macrophages and neutrophils) fibrosis in rats, elevated protein and lactate dehydrogenase levels. Bronchoalveolar hyperplasia and metaplasia, pulmonary tumorigenesis	Bermudez et al. (2004)
Ultrafine TiO₂ ∼20 nm	104 μg or 200 μg for 24 h	Intra tracheal instillation	Rats	Ultrafine particles TiO_2 increased pulmonary toxicity, as indicated by increased alveolar epithelial permeability (protein leakage), and acute pulmonary inflammation and lung injury.	Oberdörster et al. (1992)
Ultrafine TiO ₂ P25 or fine (25-300 nm)	10 mg m ⁻³ for 24 months	Inhalation exposures	Rats	Chronic inhalation exposure to 10 mg $\rm m^{-3}~TiO_2$ for 24 months causes pulmonary fibros is and lung tumours.	Heinrich et al. (1995)
TiO ₂ , rods (~35 nm) and dots (~10 nm)	1 or 5 mg single dose exposure and the recovery up to 3 months	Intra tracheal instillation	Rats	The exposure to nanoscale TiO_2 rods or dots particles produced temporary lung inflammation and fibrosis with cell injury effects at 24 h of exposure. Dots exhibit moderate to severe aggregation while rods are individually separated with little or no aggregation.	Warheit et al. (2006)
Nano sized TiO_2 (25 and 80 nm)	5 g kg ⁻¹ for 2 weeks	Oral administration	Mice	TiO_2 (25 and 80 nm) groups showed acute toxicity indicated by high coefficients of liver with hepatic injury, nephrotoxicity, myocardial damage and accumulation of TiO_2 particles in spleen, kidney and lung tissue.	Wang et al. (2007)
TiO ₂ (21 nm)	60–600 mg l ⁻¹ for 5 days	Orally via drinking water	Mice	Inflammation, oxidative stress and DNA damage in various tissues.	Trouiller et al. (2009)
TiO ₂ (50-120 nm)	5 g kg ⁻¹ for 7 days	Oral administration	Mice	Liver and kidney damage only in combination with lead acetate.	Zhang et al. (2010)
TiO ₂ (5 nm)	$5-50 \text{ mg Kg}^{-1}$ for 60 days	Intragastric administration	Mice	Brain damage morphological signs, neurobehavioral performance decline and damage of spatial recognition memory behavior.	Hu et al. (2010)

Table 1.2 Continued.

TiO ₂ NPs Information	Dose / Time	Exposure Route	Species	Toxic Effects	Author
Ultrafine TiO ₂ (21 nm) P25 75% anatase, 25% rutile.	0.1,0.5 or 1.0 mg l ⁻¹ for 14 days	Aqueous exposure	Rainbow trout (Oncorhynchus mykiss)	Mainly the anatase form: gill pathologies including oedema, thickening of the lamellae and injuries with respiratory toxicity and some irritation to the gut showing erosion of the villi tips, fusion and vacuolation of the mucosa epithelium.	Federici et al. (2007)
TiO ₂ (50 nm)	0–250 mg l ⁻¹ for 20 days	Aqueous exposure	Juvenile Common carp (<i>Cyprinus</i> <i>carpio</i>)	Exposed the fish to 100 and 200 mg l^{-1} TiO ₂ NPs for 20 days caused respiratory distress including oedema and thickening of gill lamellae, liver pathologies including necrotic and apoptosis hepatocytes and oxidative stress.	Hao et al. (2009)
TiO ₂ (24 nm)	0-100 mg Kg ⁻¹ for 8 weeks	Dietary exposure	Rainbow trout (Oncorhynchus mykiss)	50% inhibition of Na+/K+-ATPase activity seen in the brain and a 50% reduction in thiobarbituric acid reactive substances (TBARS) in the gill and intestine with Ti accumulation in gill, gut, liver, spleen and brain.	Ramsden et al. (2009)
TiO ₂ NPs 30 nm filtered / 100-500 nm unfiltered	0.2-10 mg I^{-1} for 48 h filtered / 50-500 mg I^{-1} unfiltered for 48 h	Aqueous exposure	Daphnia magna	100% mortality rate for the filtered and 9% mortality rate for the unfiltered TiO_2 NPs.	Lovern and Klaper (2006)

Table 1.3 Toxic effects of TiO₂ NPs in vitro.

TiO ₂ NPs Information	Dose / Time	Exposure Route	Toxic Effects	Author
Ultrafine TiO ₂ Anatase (10-20 nm) Rutile (200 nm)	10 μg ml ⁻¹ (about 1.77 μg cm ⁻²) for 1 h	Human Bronchial Epithelial Cells, BEAS-2B	The anatase form was much more toxic than the rutile form which caused DNA damage that detected in treatment with 10 μ g ml ⁻¹ anatase TiO ₂ 10 nm particles but no damage was detected with 5 μ g ml ⁻¹ anatase TiO ₂ 10 nm particles or with > 200 nm particles, lipid peroxidation and micronuclei formation (damage to nucleus).	Gurr et al. (2005)
Ultrafine TiO ₂ Anatase form 30-40 nm	5 μ g ml ⁻¹ suspension for 4-24 h	Human Red Blood Cells	Electron microscopy showed intracellular NPs aggregates (RBC uptake) with no morphological changes (no cellular damage).	Rothen-Rutishauser et al. (2006)
TiO ₂ NPs different sizes	From 3-30 mg ml ⁻¹ for up to 48 h	Human Lung Epithelial Cells (A549 cells)	Dose-dependent decrease in cellular viability, some lactate dehydrogenase release from cells at the highest concentration. Inflammatory mediators (interleukin 8) induced at the highest concentration.	Sayes et al. (2006)
TiO ₂ (~20nm)	$0.25-1.5 \text{ mg ml}^{-1}$	Human Fetal Lung Fibroblasts (HFL1)	Dose dependant manner toxicity effect, mitochondrial dysfunction, apoptosis and morphological modification.	Zhang et al. (2011)
$TiO_2(25 nm)$	0.8 μ g ml ⁻¹ and above for 6 h	Human Epidermal Cell Line (A431)	Demonstrate genotoxicity and cytotoxicity of epidermal cells by induces ROS and oxidative stress leading to oxidative DNA damage and micronucleus formation.	Shukla et al. (2011)
TiO ₂ P25 (30 nm)	0.2, 2 and 20 mg /cm ⁻² for 24 h	Primary Human Foreskin Keratinocytes [PHFK]	Early keratinocyte differentiation related with decrease cell proliferation and altered the calcium homeostasis.	Simon et al. (2011)
TiO ₂ NPs different sizes	10 g l ⁻¹ water suspension added to the MD exposure for 14-20 h at 36 °C	<i>Escherichia coli</i> and <i>Bacillus subtilis</i> in Minimal Davis medium(MD)	Bacterial growth inhibition (increased in presence of light). E.coli showed 72% growth inhibition in cells exposed to 5000 ppm TiO_2 , B.subtilis showed 75% growth inhibition with 1000 ppm TiO_2 and 99% growth reduction with 2000 ppm.	Adams et al. (2006)

1.6.2 Mechanisms of TiO₂ toxicity

In addition to surface area, and the chemical reactivity of particular crystal structures, there are several experiments that suggest the possible mechanisms of toxicity. TiO₂ may not be chemically modified inside organisms. For example, hydrophobic TiO₂ formulations are acutely toxic, but there is no significant difference in hydrophilic and hydrophobic functionalized TiO₂ after intratracheal instillation (Rehn et al., 2003). However, one basic toxic mechanism may involve the generation of reactive oxygen species (ROS). TiO₂ NPs have been shown to trigger ROS produced by alveolar macrophages, phagocytes and microglial cells (Olmedo et al., 2005; Long et al., 2006), whereas the ultrafine-TiO₂ (TiO₂ NPs) appears further enhance the formation of ROS in comparison to fine TiO₂ (Scherbart et al., 2011). Under ultraviolet (UV) light, TiO₂ may induce the formation of ROS such as hydrogen peroxide, hydroxyl radicals, and super oxides (Fig. 1.1). These oxygen species may lead to cellular toxicity (Blake et al., 1999) and observed DNA damage resulting in cell death after exposure to TiO_2 NPs (Trouiller et al., 2009; Shi et al., 2010). Apoptosis and cell necrosis have been also reported as cellular toxic effects of nanoscale TiO₂ which may be induced by the formation of ROS (Rahman et al., 2002; Braydich-Stolle et al., 2009; Lapied et al., 2011). This apoptosis that is induced by TiO_2 NPs has been suggested to involve the mitochondrial apoptosis pathway (Shi et al., 2010). Moreover, the phototoxic damage associated with a decrease in cell viability and an increase in the generation of ROS was also indicated after exposure of ARPE-19 cells to TiO_2 NPs under UV light (Sanders et al., 2012). Furthermore, the photocatalytic properties of TiO₂ mediating toxicity have been shown to eradicate cancer cells, and it is hypothesized that TiO_2 NPs accumulation on cell membranes may lead to cell rupture and then cell death (Thevenot et al., 2008).



Figure 1.1 Possible toxic mechanisms of TiO_2 and the generation of ROS in the cells after been exposed to UV light, showing the pathways of these oxygen species which may lead to DNA damage, membrane damage and then cell death through different cellular toxic effects.

1.6.3 Oral uptake of TiO₂ NPs and dietary exposure

Bulk TiO₂ powders have been considered safe and used as a digestibility marker (an inert marker in the diet that is not appreciably absorbed) in nutrition studies on a range of animals including fish (Mamun et al., 2007). Recently the nanoscale versions of TiO₂ have been used in personal care products (lipstick, toothpaste, sunscreens) and food (food colours, packaging, food additives E171) (Fröhlich and Roblegg, 2012). This has led to some concerns about oral toxicity of TiO₂ NPs and their uptake through the gastrointestinal tract with potential for subsequent effects on the internal organs. Powell et al., (2010) estimated the intake of TiO₂ NPs through food at about 5 mg person⁻¹ day⁻¹. Another estimation via the food by Weir et al. (2012) showed an average daily TiO₂ intake of 1-3 mg Kg⁻¹ day⁻¹, about 36% of the TiO₂ was estimated to be in the nanoscale range. Clearly there is a concern that in the long term, this daily intake may cause persistent accumulation of TiO₂ NPs in the human body.

The mechanisms that are involved in the absorption, distribution, metabolism and excretion (ADME) of the NPs remain to be investigated (Handy et al., 2008a) and they are likely to be influenced by different parameters including the aggregation, agglomeration, dispersability, size, solubility, and surface area, charge and other physico-chemistry of the NPs (Stone et al., 2010). The physico-chemical properties of NPs may play an important role in their absorption. Absorption is the first step of biological uptake for substances (Handy and Eddy, 2004). For nanomaterials, the considerations include the aggregation or precipitation of NPs during their adsorption onto the exterior surface of the organisms. There is some reliable evidence that supports this notion through experimental observation, for example, the aggregation of the TiO₂ NPs around the microvilli of the lugworm gut epithelium cells (Galloway et al., 2010). The next step is the uptake of the NPs through the cell membrane and the mechanisms of this uptake are still unclear. The possible uptake mechanisms of NPs through the gut epithelium are shown in (Fig. 1.2) which is modified from Handy et al. (2008b). The NPs are too large to be taken by ion transports across the cell membranes or by diffusion, however diffusion cannot be excluded for lipophilic NPs. Additionally, the diffusion through the paracellular route is difficult because of the ion-rich environment of the tight junctions with Ca^{2+} and Mg^{2+} which suggests that NPs would aggregate rather than diffuse. The movement of the NPs across the gut epithelium might occurby other processes like endocytotic transvesicular transport, and/or endocytosis by M-cells in the Peyer's patch (Mcclean et al., 1998; Des Rieux et al., 2007). The use of vesicular membrane trafficking systems is seemingly the most probable transport modes.

To date the information on the distribution of the NPs in the body and the target organs has yet to be established. However, the measurement of the NP concentrations in the tissue can be clarified by using inductively coupled plasma mass spectrometry (ICP-MS) (Zhang et al., 2010), and particles can also identify in the tissues using electron microscopy (Scown et al., 2009). So far there is no specific study regarding the metabolism or excretion of the NPs. However, the excretion of the NPs might occur via the process of exocytosis (Handy et al., 2008a). Simon et al. (2011) suggest that the exocytosis of TiO₂ NPs can be mediated by endosomes or other secretory vesicles.

Limited studies have been performed on the toxicity of TiO_2 NPs via oral exposure. Aqueous exposure of rainbow trout to 0 - 1.0 mg l⁻¹ TiO₂ NPs for less than 14 days caused an increase in thiobarbituric acid reactive substances (TBARS) in the intestine that suggested the fish suffered from oxidative stress (Federici et al., 2007). Some irritation was observed in the gut showing erosion of the villi tips and also fusion and vacuolation of the mucosa epithelium. The fish with severe injuries showed mucous residue in the lumen with milky colouration, which raises the possibility of TiO₂ NPs causing toxic effects via oral exposure (Federici et al., 2007). The administration of 5 g kg⁻¹ body weight of TiO₂ suspension by single oral gavage for 2 weeks in mice has

shown the ability of TiO₂ NPs (25 and 80 nm) to enter the blood circulation leading to damage in both the liver and kidneys with hepatic injury, nephrotoxicity, myocardial damage and accumulation of TiO₂ particles in spleen, kidney, lung tissue and even the brain (Wang et al., 2007). The same observation has been shown after dietary exposure (TiO₂ incorporated in food pellets) of juvenile rainbow trout to 10 or 100 mg kg⁻¹ of TiO₂ NPs for 8 weeks which resulted in Ti accumulation in the gill, gut, liver, spleen and brain (Ramsden et al., 2009). All these findings support the potential of nanoscale TiO₂ to be absorbed via the gut epithelium and subsequently pass through to other internal organs after oral administration. An *in vitro* study by Koeneman et al. (2010) provided the evidence of the ability of TiO₂ NPs to cross the human intestinal cell line (Caco-2) at a level of 10 μ g ml⁻¹ and above. However, the precise mechanism of TiO₂ NPs transfer through the gastrointestinal tract remains to be clarified. Zhu et al. (2010) also raised concerns about potential food chain effects. Their study reported the transfer of TiO₂ NPs from *Daphnia magna* to zebrafish (*Danio rerio*) through dietary exposure.

1.7 In vitro techniques for studying metal uptake

There is a strong demand for the use of *in vitro* techniques in nanotoxicology and studying the uptake mechanisms of NPs. These techniques need to be scientifically robust and ethically sound by reducing the use of laboratory animals for toxicity testing. Low-cost, high throughput *in vitro* toxicity assays should allow detailed examination under controlled conditions of various factors involved in toxicity (Panessa-Warren et al., 2006; Kahru et al., 2008). This will enhance understanding the effects of NPs at the level of physiology and aid in predicting the human health hazards.

The REACH (Registration, Evaluation, Authorisation of Chemicals) legislation aims, since it came into force on the 1st June 2007, to improve the protection of human health and the environment by the better and earlier identification of the intrinsic properties of chemical substances (Crane et al., 2008). One concern about REACH is that the novel and varied properties of NPs will generate a very large number of "new substances" that need testing. The increasing use of different NPs in a wide range of applications further highlights the need for rapid *in vitro* techniques for testing those NPs. An intelligent strategy of *in vitro* safety testing was proposed by Roszak et al. (2013) for evaluation and prediction of risk assessment of nanomaterials. This includes further steps to check cytotoxicity or genotoxicity potential of TiO₂ NPs. *In vivo* testing is a good starting point for the overall testing strategy, and could start with simple measurements such as an assessment of cell viability.

1.7.1 Perfused organ preparations

These preparations were developed to study physiology at the organ level, and have been used for the isolated perfused heart (Del Nido et al., 1998; Khairallah et al., 2004), isolated perfused liver (Strubelt, 1996; Whihelm et al., 1996), isolated perfused kidney (Frödin, 1975), isolated perfused head (gill) (Payan and Matty, 1975; Perry et al., 1984; Campbell et al., 1999) and isolated perfused intestine (Wapnir, 1991; Handy et al., 2000). Many of these isolated perfused preparations were originally used to study ion transport processes and have advantages over other techniques because the method is rapid, with the avoidance of long periods of ischemia. In addition, the volume of the external medium can be modest (e.g., 150 ml for gill perfusion, Perry et al., 1983; 500 ml for gut perfusion, Handy et al., 2000). These preparations have different layers of tissues, which mimics the situation *in vivo*, allowing the investigation of the uptake and metabolism of different materials. This may be particularly important for testing new substances like NPs. However, some difficulties are encountered with the perfusion techniques such as oedema, leakage of the perfusate and loss of the mucous constituents if the composition of the physiological saline is not carefully considered.

1.7.2 Metal uptake studies using isolated intestine preparations

The isolated gut sac preparation has been used to determine which sections of the gut are involved in metal uptake, and the perfused intestine has been used to measure the uptake kinetics and pharmacology of metal uptake (see Table 1.4). Gut sacs of oesophagus, stomach, pyloric caeca, mid and hind intestine were prepared from the whole gut and filled with the material under investigation and left for up to 4 h of exposure (Handy et al., 2000). The whole gut sac method allows a rapid way to determine in which part of the gut the metal uptake is happening. Studies on the uptake and accumulation of the metals along the gut sac showed that the main accumulation of metals was in the mid and hind intestine with 70% or more being accumulated in the mucosa rather than the underlying muscularis (e.g., copper, Handy et al., 2000; mercury, Hoyle and Handy, 2005).

The isolated perfused intestine technique has been used in both mammalian and fish physiology for many years to study the uptake of ions and water (e.g., Ando et al., 1986), and has also been used for measuring the uptake of toxic metals like copper and mercury (e.g., Handy et al., 2000). The approach involves removing the intestine from the animal, carefully turning it inside out (everted) so that mucosal epithelium is now on the "outside" and can be directly bathed in the external media (usually in a static, stirred bath of a gut saline and gassed with 95% O_2 : 5% CO_2 at room temperature, for example, 22 °C for catfish perfused intestine). This enables the experimenter to change the composition of the bath (i.e., the mucosal or luminal-side solution) so that uptake mechanisms can be investigated by adding, for instance, different concentrations of the test substance or inhibitors to the bath. Criteria for the viability of preparations are achieved by observing the normal morphology, steady appearance of the flow rate, low LDH leak and normal content of Na⁺, K⁺ and pH in the solution. Only those preparations showing normal contractions after 5-10 min resting in oxygenated saline

and with healthy morphology should be used. All these aspects (from the well preparation of the perfused intestine to good viability criteria with short time gaining results) indicate that perfusion techniques can be used effectively to study the physiology and the uptake mechanism of NPs.

		Perfused Intestine		
Chemical	Concentration	Species	Measured Factors	Author
Copper (Cu)	10, 50, 100 μmol l ⁻¹ for 2 h at 22 °C	African walking catfish (Clarias gariepinus)	Intestinal Cu uptake, effects of ion-transport inhibitors and manipulating Cl ⁻ gradient.	Handy et al. (2000)
Inorganic mercury (Hg)	0-100 μ mol l ⁻¹ for 4 h	Rainbow trout (Oncorhynchus mykiss)	Regional intestinal uptake of Hg, effects of amiloride and Ca^{2+} chelators.	Hoyle and Handy (2005)
Methyl mercury CH ₃ Hg(II)	Different concentrations at 20 or 4 °C for 1h and 45 min	Channel Catfish (Ictalurus punctatus)	Temperature sensitivity, transport mode (active/ passive process or energy dependent).	Leaner and Mason (2002)
Inorganic mercury (Hg) and methyl mercury CH ₃ Hg	50 ng ml $^{-1}$ for 2 h	Blue crab (<i>Callinectes</i> sapidus)	Accumulation and transport mode of each form of mercury in intestine and gill.	Laporte et al. (2002)
Copper (Cu) and Sodium	Cu 31.5 μ m (2 mg l ⁻¹) in isotonic solutions	Rat	Effect of Na^+ addition on Cu retention, effects of amiloride and furosemide.	Wapnir (1991)
Cu and Competitor Zinc (Zn)/ L-histidine (His)	0.1 mM Cu for 2 h	Juvenile Rats	Effect of competitors (Zn / His group) on Cu absorption, effect of high Na^+ intake and regional accumulation.	Wapnir and Lee (1993)
Inorganic mercury (Hg) and cadmium (Cd)	20 μm Hg at 37 °C for $~0.5$ or 3 min	Rat	Jejunal uptake of Hg and Cd, temperature sensitivity and mucosal permeability.	Foulkes and Bergman (1993b)

Table 1.4 Uses of the perfused intestine preparations for investigating metal uptake

1.7.3 Human colon adenocarcinoma cell line (Caco-2 cells)

The most popular intestine cell line model is the Caco-2 which was originally obtained from a human colon adenocarcinoma and has several morphological and functional characteristics of the intestinal epithelium including the presence of tight junctions in highly polarised cells (Grasset et al., 1985) and a well differentiated brush border on the apical surface (Hauri et al., 1985). The Caco-2 cell line has been used in different metal studies to assess the uptake mechanisms and toxic effects (Reeves et al., 1998; Han and Wessling-Resnick, 2002; Zerounian et al., 2003).

The use of NPs as drugs and in food has increased interest in studying the transepithelial transport behaviours of Caco-2 cells. The ability of salmon calcitonin (sCT) with biodegradable poly (lactic-co-glycolic acid) (PLGA) NPs to transport from the apical side to the basolateral side and across the monolayer of the Caco-2 cells was shown in comparison with free sCT (Sang Yoo and Gwan Park, 2004). Varied uptake capacity has been observed with different concentrations of biodegradable lactic acid – glycolic acid copolymer (PLGA 75:25) NPs (over 25-800 μ g ml⁻¹) and the maximum uptake rate occurred after 4 h exposures of Caco-2 cells (Katsikari et al., 2009). Exposure of the cells to quantum dots (QDs) coated with hydrophilic thioglycolate capping ligands at a concentration of 0.1 mg l⁻¹ induced cell death along with disruption of the epithelium monolayer (Koeneman et al., 2009). After exposure of Caco-2 cells to 10 μ g ml⁻¹ and above of TiO₂ NPs, some alteration occurred to the microvillar organization on the apical surface and an elevation of the intracellular free calcium was noticed with no cell death found or any lethal effects (Koeneman et al., 2010).

1.8 Hypothesis

The hypothesis of this study is that TiO_2 NPs can cross the epithelial layer of the gastrointestinal tract and are then taken up to the blood side using a number of active transport modes (Fig. 1.2). This idea is supported firstly by the translocation of Ti in other systemic organs, with some gut pathology indicated in freshwater fish after *in vivo* aqueous exposure (Federici et al., 2007) and dietary exposure (Ramsden et al., 2009) to TiO_2 NPs, and secondly by the indication of the ability of TiO_2 NPs to penetrate and cross the epithelial lining of *in vitro* Caco-2 cells without disrupting junctional complexes (Koeneman et al., 2010).

In this thesis, the following hypotheses are tested:

- i. Bulk TiO_2 and TiO_2 NPs will be uptake by the intestinal tissue through a form of diffusion. This could be tested by the dialysis experiment.
- ii. TiO_2 particles might be bind to the strands of mucous in the intestine so that the surface binding experiment will be applied to investigate this process.
- iii. Bulk TiO₂ and TiO₂ NPs will be uptake by the isolated perfused intestine of rainbow trout (*Oncorhynchus mykiss*) as well as by the human intestinal Caco-2 cells using an active mode of transport, either by vesicular transport of NPs or through M-cells (Fig. 1.2). Different inhibitors will be used to investigate on the uptake mechanisms mode of TiO₂.
- iv. Uptake will occur without loss of tissue integrity. The viability of the perfused intestine can be examined using standard viability markers such as LDH activity, histopathology and flow rate of the perfusion along with the leak of K⁺ and Na⁺ ions.





1.9 Aims of the research

There is a lack of information on the uptake of NPs along with the mechanisms of this uptake through the gut epithelium of different organisms. The overall aim of this study is to assess the uptake of TiO_2 NPs across the gastrointestinal tract of rainbow trout and humans using the isolated perfused intestine and Caco-2 cells, respectively. TiO_2 NPs are selected because of the wide use of these particles in the food industry and because they are incorporated in many different healthcare products which are in daily contact with humans (Aitken et al., 2006; Weir et al., 2012). Furthermore, some investigations on the physiology, behaviour and possible toxic effects will be conducted to evaluate the hazards of human exposure to TiO_2 NPs.

The specific objectives of the study are:

- i. The isolated perfused intestine and the whole gut preparation are suitable to be used in the study of nanoparticles.
- ii. Determine which part of the gut is responsible for the TiO₂ uptake using the isolated whole gut sac preparation.
- iii. Measure the uptake rate of TiO_2 across the intestinal epithelium of isolated perfused trout gut preparation.
- iv. Determine by which mechanism the uptake is happening using different inhibitors.
- v. Comparative study between different gas supplement effects on TiO₂ uptake across the intestinal epithelium.
- vi. Comparative toxicity, how the above compare to human Caco-2 cells with the consequences for hazard and risk assessment in the environment and for human health by considering the important elements of the environmental risk assessments, including the exposure and the effects (hazard) assessments.

Chapter 2

General Methodology

2.1 Stock animals

Rainbow trout, Oncorhynchus mykiss, weighting 295 ± 10 g (mean \pm S.E.M., n = 89), were collected from a commercial supplier (Torre Fisheries, Watchet, Somerset, UK). The fish were brought into the aquarium facility at Plymouth University and transported in water aerated with 100% O₂. Fish were then acclimatized in a recirculating system of approximately 6 m^3 of freshwater. Fish were kept in a photoperiod of 12 light:12 dark (on at 8 am and off at 8 pm) and held in dechlorinated Plymouth tap water with an ionic composition as following (mmol l^{-1}): Na⁺, 0.43 ± 0.01; K⁺, 0.03 ± 0.01; Ca²⁺, 0.45 ± 0.01; Mg^{2+} , 0.07 ± 0.01 (mean ± S.E.M., n = 71), and the background Ti concentrations in the water was $1 \pm 0.1 \text{ µg l}^{-1}$ (mean \pm S.E.M., n = 8). Water samples were checked daily (mean \pm S.E.M., n = 85) for pH, 6.95 \pm 0.05; temperature, 15.6 \pm 0.3 °C and oxygen saturation, $100 \pm 0.3\%$ (HACH HZ40d multi meter); and monitored weekly (mean \pm S.E.M., n = 10, in mg l⁻¹) for total ammonia, 0.12 ± 0.02 (HI95715, Hanna Instruments); total nitrite 0.06 ± 0.01 (HI93707, Hanna Instruments) and total nitrate, 3.63 ± 0.64 (HI93728, Hanna Instruments). Where necessary, a 10% trickle of new water was used to top up the recirculation system. Fish were initially fed with a food top dressed with ascorbic acid (vit C) on a normal salmonid commercial diet from EWOS. This was used in animal welfare to aid the recovery of the animals from any transport stress on arrival at the aquarium facility. Then the fish were feed twice daily with (EWOS sigma 50) which they contains: raw material (100 - 40%) inclusion), fish meal (25 - 10% inclusion), fish oil 23\%, protein 45\%, fibre 1.0%, Ash 10%, wheat, soya bean meal, beans, wheat gluten, sunflower meal, vitamins (vit A, 10 000 IU kg⁻¹; vit D, 1750 IU kg⁻¹ and vit E, 240 IU kg⁻¹), minerals, copper (Cupric sulphate 8 mg kg⁻¹) and ethoxyquin (an approved antioxidant). Before the experiments, all fish were starved for 48 h to let the gut empty of food and to facilitate the eversion of the gut.

2.2 Preparation of the isolated perfused intestine

Uptake in the preparation is generally recorded in two ways: (i) measurement of the test substance in the gut tissue at the end of the experiment, (ii) measurement of the appearance of test substance on the blood side of the preparation (i.e, in the serosal perfusate). In the latter, which represents transepithelial uptake across the gut into the blood, the perfusate is usually flows constantly in the preparation (via a peristaltic pump) to prevent the system from coming into a diffusional steady-state so that true uptake rates can be measured. In this study the isolated perfused intestine was prepared exactly according to Handy et al. (2000). The fish were humanely sacrificed by a blow to the head and the brain was pithed. The weight and total length of the fish was measured; then the entire intestinal tract (from pyloric caeca to the rectum) was quickly removed. The mass and length were also recorded for the mid and hind intestine together after cutting the tissue at the junction with the pyloric caeca. The gut was then everted, connected to the perfusion apparatus, and bathed in 500 ml of the physiological saline which contained (in mmol l^{-1}) according to Handy et al. (2000): NaCl, 117.5; KCl, 5.7; NaHCO₃, 25.0; NaH₂PO₄ 2H₂O, 1.2; CaCl₂ 2H₂O, 5.0; MgSO₄, 1.0; glucose, 5.0; mannitol, 23.0; osmolarity ~320 mosm (Osmomat 030, Gonotec) and adjusted to pH 7.4 (Hanna GLP Bench-top pH/mV/ISE/°C meter) with a few drops of 1 mol l⁻¹ HCl, at 18°C (a constant room temperature), gassed with 95% O₂: 5% CO₂ (a standard gas mixture used for gut perfusion work, Handy et al., 2000) or with 99.5% O₂: 0.5% CO_2 ("low CO_2 " gas mixture). The latter, being closer to the acid-base status of trout in vivo (Milligan et al., 1991; Goss et al., 1992). The intestine was serosally perfused (1 ml min⁻¹ via an Ismatec peristaltic pump) using the gut saline described above for up to 4 h (Fig. 2.1). This physiological saline is a buffered salt solution which is used for the both sides of the tissue (symmetrical perfusion) and is sufficient to meet the ionoregulatory and acid-base needs of the tissue. Perfusions were conducted in pairs, with a control (no added TiO₂) and treatment (e.g., + TiO₂ NP or bulk powder) performed each day (n = 6 perfusions per treatment; n = 7 perfusions for the control and TiO₂ NP treatment gassed with the standard gas mix). The eluted serosal perfusate was collected manually in 10 min fractions, and the volume determined gravimetrically. Five ml of the mucosal solution (bath samples) were collected every 30 min for TiO₂ analysis (see below), and the pH was checked. At the end of each experiment the gut was carefully washed in deionised water to remove excess TiO₂, and the surface area of the open gut was measured (manually using graph paper). Samples of each gut were taken for histopathology and electron microscopy. Then the gut was cut into mid and hind sections for trace metal analysis and moisture content (see below), while the perfusate and mucosal solutions were analysed for TiO₂ and biochemistry (see below) (Fig. 2.2).



Figure 2.1 Diagram (A) showing the processing and the design of the isolated gut perfusion experiment. The images in (B and C) presents the everted mid and hind intestine bathed into 500 ml of the physiological saline, gassed with 95% O_2 : 5% CO_2 or 99.5% O_2 : 0.5% CO_2 and serosally perfused with another volume of physiological saline for up to 4 h.



Figure 2.2 Experimental processing of the isolated perfused intestine after a 4 h perfusion. Mid and hind intestine were digested in a concentrated nitric acid and then analysed by the ICP-OES for the Ti and metal concentrations in the tissue. LDH assay was applied to investigate on any tissue damage by testing the perfusate and mucosal (bath) solutions and then those solutions were analysed by the ICP-OES for the Ti and metal concentrations. Distributions of the TiO₂ particles in the mucosal solution were further detected using the nanosight. A piece of intestinal tissue were taken from the middle part of the gut after 4 h perfusion for histopathology and a small pieces also taken from the same area for TEM examination.

2.3 Lactate dehydrogenase (LDH) assay

The routine viability criteria for the perfused intestine preparation are the presence of steady perfusate flow, net efflux of water across the gut in a freshwater fish (see Handy et al., 2000), as well as the LDH leak measurements. In the perfused intestine LDH leak from the tissue is usually low and negligible (Handy et al., 2000) and any elevation of this cytosolic enzyme can indicate tissue damage to the preparation. LDH measurements were done, along with perfusate flow and net water flux, to determine if these criteria also worked for perfusions with nanomaterials. After each experiment all the samples (perfusate and mucosal solutions) were analysed for the LDH activity. The LDH assay was determined by the disappearance of nicotine amide adenine dinucleotide (NADH) at 340 nm (Helios B, Spectrophotometer) resulting from NADH oxidation by pyruvate at 25°C and pH 7.4. According to Campbell et al. (1999), the reaction mixture contained: (i) 2.8 ml of 0.6 mmol l^{-1} pyruvate in 50 mmol l^{-1} phosphate buffers (pH 7.4); (ii) 0.1 ml of 0.6 mmol l⁻¹ NADH solution; (iii) 0.1 ml of sample then mixed directly in a 3 ml cuvette. The reaction was allowed to proceed for 2 min and the LDH concentration calculated using an extinction coefficient of 6.3 for a path length of 10 mm. LDH activity is expressed as IU ml⁻¹ (equal to μ mol min⁻¹ ml⁻¹) for individual samples and as IU (µmol min⁻¹) for the cumulative activity of the LDH. The samples and blanks were compared with an LDH standard (0.1 ml of 10.66 IU ml⁻¹). Typically, in perfused organ preparations from fish, the maximum LDH release into the media from a normal healthy tissue would not exceed 1 IU ml⁻¹ and the data for all samples here were below this value (see results).

2.4 Trace metal analysis

The methodology for trace metal analysis in tissues was based on the original nitric acid digestion method of Handy et al. (2000) for fish intestine, but with some novel

modifications that enable detection of Ti from intact TiO₂ NPs dispersed in the tissue digest without the need for aggressive hydrofluoric acid or hot sulphuric acid digestion (see Lomer et al., 2000 for the traditional H₂SO₄ method for foods) to completely dissolve the Ti metal (Ramsden et al., 2012 in review, and patent pending, UK Patent application No: 1207745.9). The modification presented here involved adding Triton X-100 to a diluted the nitric acid digest, and with mixing/shaking it was possible to get good nebulisation of the sample in the spray chamber of the inductively coupled plasma optical emission spectrometer (ICP-OES, Varian 725 ES) to detect both the total Ti metal concentrations from the NP exposure and other dissolved trace metals in the sample. Briefly, tissues from all experiments were oven dried to a constant weight at 100 °C for up to 48 h (Gallenkamp Oven BS Model OV- 160), then the samples were transferred into 20 ml scintillation vials (VWR International Ltd, Poole, UK) and digested in 1 or 4 ml (depending on tissue weight) of concentrated nitric acid for 3 h at 70 °C in a water bath. The digested samples were allowed to cool, and then Triton X-100 was slowly added to each digested with Milli-Q water (ultra-pure ion free water) to achieve a final volume of 2% Triton X-100 in each sample. Typically this involved carefully adding around 3.2 or 0.8 ml of 10% Triton X-100 and diluting to a final volume of 16 or 4 ml (for tissue weights of $\sim 1g$ or 0.1g respectively). Digests were then analysed for total Ti metal, Na⁺, K⁺, Ca²⁺ and Mg²⁺ concentrations by ICP-OES. Critically, samples were mixed immediately before introduction to the instrument (30 min, KS501 digital orbital shaker, IKA Labortechnik; set at 145 r min⁻¹, or vortexing for 10 s IKA MS2 Minishaker; set at 2500 r min⁻¹), depending on the size/shape of the test tubes used. The instrument detection limit (3 x standard deviation of the blank) for measuring the gut salines or matrix matched standards was calculated as 1.920 μ g l⁻¹ (equivalent to 0.04 µmol l⁻¹ of Ti metal), and for the entire digestion protocol (procedural blank) was 4.925 μ g l⁻¹ (equivalent to 0.1 μ mol l⁻¹ of Ti metal, and for a

typical 0.5 g tissue sample equates to 0.0032 μ mol Ti metal g⁻¹ dry weight of tissue). The optimised protocol gave good spike recovery for fish tissue (e.g., mean \pm SEM, n = 6-13; 97.5 $\pm 2.9\%$ for spiking with Ti metal, and 92.2 $\pm 1.4\%$ for spiking with TiO₂ NPs) and low coefficients of variation within and between tissue samples from different stock fish (around 5% or less for repeats of the entire protocol). Calibrations with different concentration of Triton-X100 in standards had a negligible effect (no problems with instrument calibration with metals standards or TiO₂ dispersions, Ramsden et al., in review).

Tissue Ti levels are reported as μ mol of Ti metal g⁻¹ dry weight, being converted to molar units in order to allow comparison with gut perfusion studies in the epithelial physiology literature and previous Ti measurements in trout tissue (see Federici et al., 2007). The bath solution was measured for Ti to confirm the TiO₂ dosing and for convenience the data are reported as mg l⁻¹ of TiO₂ (for the bath solution only) after correcting the Ti metal data using the stoichiometry of the compound (Ti content of TiO₂ as 60% of the mass). All other data are reported as total Ti metal concentrations (in the tissues and eluted perfusates). The moisture content of tissues was determined by taking the weight of the tissue before and after oven drying and calculated as: ((wet weight – dry weight)/wet weight) x 100. The mucosal solutions from all experiments were also analysed using a flame photometer (Corning 480) for Na⁺ and K⁺ concentrations. The latter instrument burns the sample in an air/acetylene flame, and detects changes in absorbance due to the colour intensity of each element (Domingo and klyne, 1949). The absorbances are converted to metal concentrations by the software in the instrument using standards and the application of the Beer-Lambert law.

2.5 Preparation of titanium dioxide NP and bulk stock solutions

A 1 g 1^{-1} dispersion of bulk TiO₂ was prepared from 1 g of TiO₂ powder (ACROS, Titanium (IV) oxide, New Jersey, USA; manufactures information: purity of 98.0-

100.5% TiO₂). The manufacturer's technical support indicated a crystal structure mixture of more anatase than rutile, but could not report the exact proportions. Counting of crystal shapes on our electron microscopy images (n = 10 random pictures) indicated 25.3% rutile and 74.7% anatase. The powder was dissolved and stirred (Stuart magnetic stirrer, set at half speed) in 1 litre of ultrapure Milli-Q for 20 minutes. The particle size was determined by two independent methods. First, sub samples of the 1 g l^{-1} stock dispersion in ultrapure Milli-Q water (stirred for 20 min) were examined using transmission electron microscopy (TEM, JEOL-JEM.1400) and examples of the particles are shown in (Fig. 2.3A) with a primary particle size measurement of 147 ± 9 nm (mean \pm S.E.M., n = 148), while the aggregates viewed by electron microscopy were typically 1124 ± 331 nm (mean \pm S.E.M., n = 9). The second approach for particle sizing was nanoparticle tracking analysis (NTA, using a Nanosight LM 10, Nanosight, Salisbury, UK, laser output set at 30 mW at 640 nm). In order to measure particle size distribution by NTA the initial stock dispersion $(1 \text{ g } 1^{-1})$ was diluted to 1 mg 1^{-1} in the physiological saline used for gut perfusions (see below) to account for matrix effects on aggregation (0.5 ml of the initial stock into 500 ml of saline, stirred as above), and the resulting dispersion gave a mean values of (hydrated radius) 166 ± 20 nm for the average particle size and 84.7 ± 19.4 nm mean of the smallest size bin of the particles (mean \pm S.E.M., n = 3) (Fig. 2.3C). The mucosal samples from the end of the experiments using 1 mg l^{-1} bulk TiO₂ were also analysed for Ti distribution using the Nanosight LM10 (see results, Fig. 2.3E).

The stock material of TiO_2 NP used here was the same as that used by Federici et al. (2007). The powder form of ultrafine titanium dioxide NP type "Aeroxide" P25 (DeGussa AG, supplied by Lawrence Industries, Tamworth, UK) with (revised manufactures information): a crystal structure of 25% rutile and 75% anatase TiO₂, purity was at least 99% TiO₂ (maximum impurity stated was 1% Si), the average

particle size was 21 nm with a specific surface area of $50 \pm 15 \text{ m}^2 \text{ g}^{-1}$. Chemical analysis of stock dispersions revealed no metal impurities (below the detection limit so the data are not shown), and the batch purity was high. A stock dispersion of 10 g l^{-1} TiO₂ NPs was made (no solvents) by dispersing the NPs in ultrapure Milli-Q water with sonication in a bath-type sonicator (500 ml of stock, 35 kHz frequency, Fisher brand FB 11010, Germany) for 6 h. A 1 g l^{-1} TiO₂ NPs stock solution was prepared from the stock above by diluting 10 ml to 100 ml in ultrapure Milli-Q water, and sonicated as above for 6 h. This second stock dispersion was used for dosing the perfusion bath. Transmission electron microscopy showed the structure and the measured primary particle size of 22.8 \pm 0.6 nm (mean \pm S.E.M., n = 169), while the mean size of aggregation of NPs were 121.7 ± 19.7 nm (mean \pm S.E.M., n = 8; Fig. 2.3B) with similar crystals showing as the bulk TiO₂. These measurements were very similar to that reports for TiO₂ NPs in Federici et al. (2007). The dosing of the bath was as above for the bulk material, and the distribution of the 1 mg l^{-1} TiO₂ NPs in the physiological saline prior to the start of experiments was also confirmed by NTA. The mean values for particle size were 117 \pm 17 nm and the smallest size bin of particles had a mean of 35 ± 5 nm (mean \pm S.E.M., n = 3; Fig. 2.3D). The mucosal samples from the end of the experiments following expose to 1 mg l^{-1} TiO₂ NP were also analysed for Ti concentration using the Nanosight (Fig. 2.3F).

Chapter 2



Figure 2.3 Electron micrographs showing (A) bulk and (B) nano TiO_2 particles in a 1 g Γ^1 stock dispersion. Particle size distributions measured by nanoparticle tracking analysis (NTA) are shown below; (C) bulk TiO_2 and (D) TiO_2 NPs in a 1 mg Γ^1 stock dispersion in physiological saline (without the gut present); (E) bulk TiO_2 and (F) TiO_2 NP in the same mucosal solution samples at the end of a 4 h intestinal perfusion, graphs are individual examples from triplicate measurements of each.

2.6 Histology and transmission electron microscopy

After each perfusion a piece of tissue from the middle of each preparation (so as to avoid the ends attached to the apparatus) was carefully collected for light microscopy (n = 6 perfusions per treatment; n = 7 perfusions for the control and TiO₂ NP treatment gassed with the standard gas mix). Histology was performed according to (Handy et al., 2002b). Briefly, tissues were rinsed and fixed in 10% buffered formal saline for at least 48 h, dehydrated in graded ethanol solutions to remove excess water, and cleared in three graded series of xylene for 1 h each prior to embedding in paraffin blocks. Dewaxed sections (8 µm) were stained with haematoxylin and eosin. Slides were examined using light microscopy (Olymus Vanox - TAH2) and photographs were taken using a digital camera (Olympus camedia C-2020 Z).

For the transmission electron microscope investigations, small fragments from mid and hind intestine (n = 3 per treatment) were fixed in glutaraldehyde (2.5%) at low temperatures (4 °C), buffered in 0.1 M sodium caodylate pH 7.2 and post-fixed in osmium tetra oxide 2% (OsO4). The pieces were then dehydrated in a graded series of ethanol and infiltrated with a graded series of Spur's resin. Embedding as a small capsule was performed using pure resin. Thin sections were cut and stained with 2% uranyl-acetate and lead citrate (Au et al., 1999), then examined by transmission electron microscope (TEM, JEOL-JEM.1400).

2.7 Calculations and terminology

All calculations on uptake rates were performed according to Handy et al. (2000). Briefly, data were graphically presented as the absolute cumulative appearance of Ti metal in the perfusate, plotted against exposure time for example perfusions; and with cumulative perfusate flow shown on the same graph. The absolute Ti content of each 10 min fraction (μ g) was calculated by multiplying the total Ti metal concentration in each
tube (µg ml⁻¹) by the total volume of perfusate/tube (ml). Ti contents of each fraction were summed for the cumulative plots (reported as nmols absolute of Ti metal). The initial (first 10 min) and overall (240 min) net Ti flux rates ($J_{net,Ti}$, in nmol g⁻¹ dry tissue h^{-1}) to the serosal perfusate were calculated from the perfusate Ti content (in nmol) divided by the tissue dry mass and corrected to 1 h, while the initial (first 10 min) and overall (240 min) net water flux rates (J_{net} , H_{20} , in ml g⁻¹ dry tissue h⁻¹) to the eluted perfusate were calculated from the differences between the rates of cumulative perfusate and effluent flow divided by the tissue dry mass and corrected to 1 h. In the present study, precise terminology were used for the absolute amounts of Ti metal in the perfusates (nmols, not a concentration), the concentration of total Ti metal in the tissue (μ mol g⁻¹ dry weight of Ti metal, not TiO₂ compound), and distinguish this from the mg 1^{-1} of TiO₂ compound added to the bath when confirming the exposure. The phrase "total Ti metal concentrations" is used to mean the total mass concentration of Ti (not TiO₂ compound) in the tissue or relevant salines determined by ICP-OES, it does not infer anything about whether the Ti is present as particulate TiO₂ or as a dissolved Ti species. The term "Ti accumulation" is used to mean a net increase in the total Ti metal concentration in the tissue over time, determined by ICP-OES of the tissue digests. The phrase "Ti uptake" is used specifically to mean the net uptake of total Ti metal to the serosal compartment in relation to the flux calculations above. Again, this relates to ICP-OES measurements of Ti in the perfusates and does not infer anything about whether the Ti is taken up as particulate TiO₂ or as a dissolved Ti species. In relation to the dialysis experiments, the phrase "apparent total dissolved Ti" is sometimes used to mean the total Ti metal concentration in the aqueous phase of the beaker that has passed through the dialysis membrane. Dispersed TiO₂ particles are therefore distinguished from the apparent total dissolved Ti metal in the text.

2.8 Statistical analysis

All data were presented as mean \pm S.E.M (standard error of mean) and analysed using Stat Graphics Plus Version 5.1 while figures were drawn using Excel or sigmaplot 12.0. The data (treatment and time effects) were tested after checking for kurtosis, skewedness, and unequal variance (Bartlett's test). Parametric data were tested by oneway ANOVA followed by the least squares difference multiple range test. Nonparametric data that could not be transformed were tested using the Kruskal–Wallis test (analysis by ranks) and differences were located using notched box and whisker plots. The Student's *t*-test was used as well to investigate the differences between pairs of data, or the Mann-Whitney U test where appropriate for non-parametric data. All statistical analysis used a 95% confidence limit, so that *P*-values equal to or greater than 0.05 were not considered statistically significant.

Chapter 3

Uptake of titanium from TiO₂ nanoparticle exposure in the isolated perfused intestine of rainbow trout, (*Oncorhynchus mykiss*) with novel CO₂ components

Abstract

In vivo studies have raised concerns that titanium dioxide nanoparticles (TiO₂ NPs) may be taken up across the gut of fish, but there are limited data on the uptake kinetics. Whole gut sacs and the isolated perfused intestine of rainbow trout (Oncorhynchus *mykiss*) were used to determine the parts of the gut involved in absorption, and the intestinal uptake rates of Ti from TiO₂ NPs across fish gut. Differences between bulk and nanoscale TiO₂, and the effects of varying the CO₂ content of the perfustae were also examined. Luminal exposure of whole gut sacs to 1 mg l⁻¹ TiO₂ NP for 4 h caused accumulation of Ti in the hind and mid intestine, with 80% or more of the Ti in the mucosa, rather than underlying muscularis. Perfused intestines exposure to 1 mg l^{-1} TiO₂ NP or the equivalent bulk powder in the mucosal solution for 4 h gassed with 95% $O_2:5\%$ CO₂ showed a time-dependent accumulation of Ti in the serosal perfusate with a maximum initial uptake rate (mean \pm S.E.M., n = 6 for the bulk TiO₂; n = 7 for the control and TiO₂ NP treatment) of 1.55 ± 0.33 , 0.98 ± 0.47 , and 0.23 ± 0.16 nmol g⁻¹ h⁻¹ for TiO₂ NPs, bulk powder, and no-added Ti controls, respectively (statistically significant differences on all treatments, ANOVA or Kruskall-Wallis test, P < 0.05) demonstrating NP translocation across the gut into the blood side. Notably, there was at least a 10 fold increase in Ti from TiO₂ NPs uptake to 21.16 ± 18.9 nmol g⁻¹ h⁻¹ (mean \pm S.E.M., n = 6) when the CO₂ content of the perfusate gas was reduced to 0.5% (statistically significant, t-test, P < 0.05). In conclusion, the TiO₂ NPs are mostly absorbed across the mid and hind intestine, the uptake rate for Ti from TiO₂ NPs across the intestine being much faster than the bulk powder, and the uptake mechanism(s) involve a CO₂ sensitive component across the mucosal membrane.

3.1 Introduction

The wide applications of TiO₂ NPs in food and different consumer products arise from the unique physico-chemical properties of TiO₂ at the nanoscale including its stability, photocatalytic properties, and bright whiteness as a pigment (e.g., Allen et al., 2002; Krischok et al., 2002). The respiratory toxicity of TiO₂ NPs is relatively well established in mammals (Bermudez et al., 2004; Warheit et al., 2006) and waterborne exposure of fish gills to metallic NPs also causes pathology of the respiratory epithelium (TiO₂ NPs, Federici et al., 2007; Cu NPs, Griffitt et al., 2009).

For exposure via the gut, fish that drink water containing 1 mg l^{-1} TiO₂ NP show irritation to the gut mucosa with erosion of the villi tips, fusion and vacuole formation in the mucosal epithelium (Federici et al., 2007). However, dietary exposures to TiO₂ NPs in the food are much less toxic, and trout readily eat diets containing 10 or 100 mg kg^{-1} of TiO₂ NPs for 8 weeks with a normal growth rate and haematology (Ramsden et al., 2009). The latter study also showed Ti accumulation in some of the internal organs, suggesting dietary uptake across the intestinal epithelium of either Ti metal or TiO_2 NPs. This internal uptake suggests that TiO_2 NPs do cross the gut. The uptake of TiO_2 NP across gills and gut epithelium had also been suggested from the appearance of metal from metallic NPs in waterborne studies (Federici et al., 2007; Griffitt et al., 2008; Hao et al., 2009), supports the notion of possible NP transfer across gill epithelia with the ability of NPs to cross cell membranes (Geiser et al., 2005) and penetrate into and through the cells without disrupting junctional complexes (Koeneman et al., 2010). Metallic NPs have also been located on or in gill epithelial cells of trout using coherent anti-Stokes Raman scattering (CARS) (Johnston et al., 2010). However, precise information on the rates of uptake of TiO₂ NPs across epithelia is lacking, especially for the intestine.

The aim of current study was to determine which part of the gut was involved in the Ti accumulation from an exposure to TiO_2 NPs by using isolated whole gut sacs (Hoyle and Handy, 2005), and then to measure the total Ti uptake rates based on total metal concentrations in perfusates using the isolated perfused intestine (Handy et al., 2000). These long-established *in vitro* preparations of intact intestinal tissue allow easy manipulations of the external media to measure concentration-dependent uptake. An additional aim was also to demonstrate the utility of the perfused intestine preparation for studies with manufactured NPs, and to show that viability criteria and other measurements relating to physiological function could be met. Finally, the work also aimed to explore changes of the composition of the media for NP studies by using different gas mixtures to reveal a CO_2 -sensitive component to metal uptake from particle exposure that has not been previously reported.

3.2 Methodology

3.2.1 Preparation of the whole gut sacs

Whole gut sacs were used to determine which parts of the gastrointestinal tract are mainly involved in Ti absorption from TiO_2 exposure and prepared according to Hoyle and Handy (2005) derived from the original method of Handy et al. (2000). This well-established physiological technique involves removing the whole gut from the animal, filling the lumen (according to the size of the gut trying not tighten the pressure inside the gut) with the test substance of interest, and then suturing closed the different anatomical parts of the gut so that regional uptake into the tissue can be measured over a few hours (Fig. 3.1). Fish were humanely sacrificed by a blow to the head and the brain was pithed. The weight and total length of the fish was measured; then the entire gastrointestinal tract was carefully removed, washed and flushed through with the physiological saline (described below). The gut was sutured closed at the posterior end,

and the lumen filled with 1 mg l^{-1} TiO₂ (NP, Bulk, or no added TiO₂ control as appropriate) in physiological saline which contained (in mmol 1⁻¹): NaCl, 117.5; KCl, 5.7; NaHCO₃, 25.0; NaH₂PO₄ 2H₂O, 1.2; CaCl₂ 2H₂O, 5.0; MgSO₄, 1.0; glucose, 5.0; mannitol, 23.0; osmolarity ~320 mOsm (Osmomat 030, Gonotec) and adjusted to pH 7.4 with a few drops of 1 mol l^{-1} HCl, at 18 °C, gassed with 95% O₂: 5% CO₂. The concentration of 1 mg l^{-1} TiO₂ was chosen because it may be physiologically relevant and to compare it with previous work on TiO_2 by Federici et al. (2007). Then the gut was closed with a suture around the upper end of the oesophagus, and further sutures applied to separate each region of the gut (oesophagus, stomach, pyloric caeca, mid and hind intestine) according to Hoyle and Handy (2005). The gut was then bathed with 500 ml of the same physiological saline (no added TiO₂). The bath (serosal side) was continuously stirred and gassed with 95% O₂: 5% CO₂ at 18 °C for 4 h. The 4 h duration time for this experiment was chosen following the previous metal perfusion studies which indicated saturable uptake within 4 h (e.g. Cu in catfish gut, Handy et al., 2000; Hg in trout gut, Hoyle and Handy, 2005). After each experiment the gut was carefully opened and washed with deionised water to remove excess TiO₂, then divided into sections of intact oesophagus, stomach, pyloric caeca, mid and hind gut for metal analysis and histology (see below). In some experiments, each part of the gut was further divided longitudinally into two equal parts; one part was stripped of the mucosa, while the other part was left whole, to enable calculation of the relative proportion of Ti in the mucosa and underlying muscularis (Hoyle and Handy 2005).

A full description of the isolated perfused intestine, LDH assay, trace metal analysis, and the stock TiO_2 solution preparations, stock animals, histology with TEM, calculations and statistical analysis were found and explained in details in Chapter 2.



Gassed & Stirred Bath



Figure 3.1 An ideal diagram of the whole gut sac preparation technique (A). The images (B and C) shows the different parts of the gut filled with the exposure material, sutured close each part separately and bathed into 500 ml of physiological saline, continuously stirred and gassed with 95% O_2 : 5% CO_2 for up to 4 h incubation.

3.3 Results

3.3.1 Ti accumulation by whole gut sacs experiment

The whole gut sac preparation was used to determine which regions of the gut were accumulating Ti from TiO_2 exposures (Table 3.1). All regions of the intact gut showed statistically significant elevations of total Ti metal concentrations in the tissues from exposure to the bulk TiO₂ compared to the unexposed control, with the greatest increases in the mid and hind intestine (Kruskal-Wallis test, P = 0.009 and 0.006 for mid and hind intestine respectively, Table 3.1). Elevations of total Ti metal concentrations in the tissues were also observed for nanoscale TiO₂ exposures in all regions of the gut, and unlike the bulk material there were no statistically significant differences between the regions of the gut in the TiO₂ NP treatment (ANOVA, P = 0.25, Table 3.1). Within each region of the gut there was no material-type effect (comparison of bulk versus nano treatment, ANOVA or Kruskal-Wallis test, P > 0.05) on the tissue total Ti metal concentration. Comparison of the proportions of total Ti metal concentrations in the mucosa and underlying muscularis of stripped gut confirmed that most (74% or more) of the background Ti in the gut of control animals was in the mucosa. Similar observations were made for guts exposed to TiO_2 NPs and bulk TiO_2 (Table 3.2). However, for bulk TiO_2 the proportion of Ti in the mucosa of the mid intestine was only 54% and even less for the hind intestine; both gut regions being much lower than the TiO₂ NP treatment which showed 71% or more of the tissue total Ti metal concentration in the mucosa.

3.3.2 Electrolyte composition and moisture content of whole gut sac experiments

There were some regional differences in the electrolyte composition of the gut as expected (Table 3.1). However, there were no treatment-dependent effects of either bulk or nanoscale TiO_2 on the Na⁺, K⁺, Ca²⁺, or Mg²⁺ concentrations in the whole gut sacs. A

slight increase in the concentrations of K⁺ and Na⁺ was detected in some regions of the gut but this increase was not significantly different (ANOVA or Kruskal-Wallis test, P > 0.05) compared to the control values. A depletion in the concentrations of Ca²⁺ and Mg²⁺ was founded in all regions of the gut exposed to TiO₂ (NP or bulk) compared to the controls; with a statistically significant difference found only in the mid intestine (ANOVA or Kruskal-Wallis test, P < 0.05, Table 3.1). An increase in the moisture content was found in all region of the gut except for the oesophagus compared to the controls after exposure to bulk or TiO₂ NPs with a statistically significant difference found in the mid intestine (ANOVA or Kruskal-Wallis test, P < 0.05, Table 3.1).

Potassium concentration in the stripped (muscularis) was higher than that found in the mucosa (Table 3.3) in all gut regions, except for the oesophagus which showed lower values of K⁺ than the mucosa. Sodium concentration in mucosa was mainly higher than the stripped one after exposed to TiO₂ NPs (66% or more) indicated by a significantly difference (ANOVA or Kruskal-Wallis test, P < 0.05) compared to the control or bulk treatment in mid and hind intestine (Table 3.3). Calcium and magnesium concentrations were accumulated in the mucosa more than the stripped gut after exposure to TiO₂ NPs for all regions of the gut; with values for the magnesium being significantly higher than either the control or bulk in the mucosa of mid and hind intestine (ANOVA or Kruskal-Wallis test, P < 0.05, Table 3.3).

3.3.3 Histopathology of the whole gut sac experiment

Morphological examination of the whole gut sac after 4 h incubation with $1 \text{ mg } 1^{-1} \text{ TiO}_2$ NPs or bulk resulted in normal gut morphology with no pathologies detected in gut tissue. However, a minor effect was observed in the pyloric caeca represented by the appearance of vacuoles in just one out of six fish examined from one treatment (Fig. 3.2-i).



Figure 3.2 Histology of the whole gut sac after 4 h experiment. (a) control oesophagus, (b) 1 mg 1^{-1} bulk TiO₂ oesophagus, (c) 1 mg 1^{-1} TiO₂ NP oesophagus showing normal tissue. (d) control stomach, (e) 1 mg 1^{-1} bulk TiO₂ stomach, (f) 1 mg 1^{-1} TiO₂ NP stomach showing normal tissue. (g) control pyloric caeca, (h) 1 mg 1^{-1} bulk TiO₂ pyloric caeca, (i) 1 mg 1^{-1} TiO₂ NP pyloric caeca showing appearance of vacuoles (V). (j) control intestine, (k) 1 mg 1^{-1} bulk TiO₂ intestine, (l) 1 mg 1^{-1} TiO₂ NP intestine showing normal tissue. Scale bar = 100 µm, sections were 8 µm thickness and stained with haematoxylin and eosin (n = 6 fish per treatment).

	[Metal] µmol g ⁻¹ dry mass									
Paramet	er	Oesophagus	Stomach	Pyloric caeca	Mid intestine	Hind intestine				
[Ti]		-								
	Control	$< 0.003^{a}$	0.003 ± 0.001^{a}	$< 0.003^{a}$	0.004 ± 0.001^{b}	0.006 ± 0.004^{a}				
	Bulk TiO ₂	$0.033 \pm 0.009^{ab,*}$	$0.029 \pm 0.007^{a,*}$	$0.016 \pm 0.003^{a,*}$	$0.062\pm 0.018^{b,\ast}$	$0.041 \pm 0.011^{ab,*}$				
	TiO ₂ NP	$0.027 \pm 0.009^{a,*}$	0.017 ± 0.003^{a}	$0.015 \pm 0.003^{a,*}$	$0.027 \pm 0.005^{a,*}$	$0.030 \pm 0.006^{\mathrm{a},*}$				
$[K^+]$	-									
	Control	$158.4\pm16.8^{\rm ab}$	168.4 ± 23.2^{a}	$87.5\pm7.9^{\rm b}$	$224.5 \pm 35.1^{\rm a}$	$274.6 \pm 71.4^{\rm a}$				
	Bulk TiO ₂	$126.7\pm18.8^{\rm ab}$	$175.8 \pm 13.7^{\mathrm{b}}$	$81.4 \pm 10.4^{\rm a}$	$248.7 \pm 15.7^{\circ}$	$239.4 \pm 29.3^{\circ}$				
	TiO ₂ NP	$221.5\pm86.3^{\rm a}$	$184.7 \pm 22.2^{ m ad}$	$80.5\pm5.8^{\rm b}$	203.1 ± 25.9^{cd}	$239.3 \pm 11.1^{\circ}$				
$[Na^+]$										
	Control	$245.9 \pm 21.6^{\rm a}$	$219.6 \pm 27.1^{\mathrm{a}}$	$121.8 \pm 16.8^{\rm b}$	$245.3 \pm 44.7^{\rm a}$	$286.4 \pm 54.7^{\mathrm{a}}$				
	Bulk TiO ₂	$207.8 \pm 25.9^{\rm a}$	$253.3 \pm 12.7^{\mathrm{a}}$	$114.9 \pm 9.1^{\rm b}$	$353.6 \pm 75.3^{\rm a}$	$290.3 \pm 53.0^{\rm a}$				
	TiO ₂ NP	$311.2\pm114.8^{\rm a}$	$207.5 \pm 23.2^{\mathrm{ac}}$	102.1 ± 7.1^{b}	$269.7 \pm 38.1^{\circ}$	305.9 ± 58.9^{ac}				
$[Ca^{2+}]$										
	Control	$17.2\pm2.2^{\mathrm{a}}$	$16.8 \pm 2.9^{\rm a}$	$10.9\pm2.8^{\mathrm{b}}$	$22.1\pm8.5^{\rm ab}$	23.9 ± 9.2^{ab}				
	Bulk TiO ₂	$12.6 \pm 1.7^{\mathrm{ab}}$	$15.6 \pm 1.3^{\rm b}$	$8.9\pm1.5^{\rm a}$	$21.6 \pm 2.9^{c,*}$	$15.9 \pm 1.4^{\mathrm{b}}$				
	TiO ₂ NP	$18.8 \pm 7.4^{ m a,*}$	$12.7\pm1.7^{\mathrm{a}}$	$6.2\pm0.5^{\mathrm{b}}$	$14.9 \pm 1.8^{ m ac,\#}$	$17.9 \pm 2.4^{\circ}$				
$[Mg^{2+}]$										
	Control	15.9 ± 1.3^{a}	$15.4\pm1.7^{\mathrm{a}}$	$9.5\pm0.8^{ m b}$	18.6 ± 3.2^{a}	22.3 ± 5.1^{a}				
	Bulk TiO ₂	$13.0\pm1.7^{\mathrm{a}}$	$16.3\pm0.5^{\mathrm{ad}}$	$8.7\pm0.9^{ m b}$	$20.9 \pm 1.5^{\circ}$	$18.8\pm2.0^{ m cd}$				
	TiO ₂ NP	$22.4\pm8.7^{\rm a}$	$15.6 \pm 1.4^{\rm ac}$	$8.4\pm0.4^{\rm b}$	$17.4 \pm 1.9^{\mathrm{ac}}$	19.5 ± 0.7^{c}				
Moisture	e (%)									
	Control	80.2 ± 2.6^{a}	$77.5\pm1.8^{\rm a}$	62.6 ± 7.2^{b}	$74.5\pm1.8^{\rm a}$	$78.2\pm2.9^{\mathrm{a}}$				
	Bulk TiO ₂	75.9 ± 0.8^{ab}	$79.6\pm0.7^{\rm a}$	68.6 ± 6.2^{b}	$83.1 \pm 1.4^{a,*}$	$80.1 \pm 1.3^{\mathrm{a}}$				
	TiO ₂ NP	76.1 ± 0.5^{a}	$78.7\pm0.8^{\rm a}$	66.3 ± 5.8^{b}	$78.4 \pm 2.2^{a,*}$	$81.7\pm0.7^{\rm a}$				

Table 3.1 Total Ti metal, K^+ , Na^+ , Ca^{2+} and Mg^{2+} concentrations of gut tissue segments from whole gut sacs following exposure to 1 mg l⁻¹ of TiO₂ in the gut lumen for 4 h.

Values are means \pm S.E.M. (n = 6 for each group) expressed as μ mol g⁻¹ dry mass of intestinal tissue, except moisture content (%) = ((wet weight – dry weight)/wet weight) x 100. Whole gut sacs were filled with physiological saline (control), 1 mg l⁻¹ bulk TiO₂ and 1 mg l⁻¹ TiO₂ NP at laboratory temperature (18°C) for 4h. * Statistically significant difference from control values within columns and electrolytes (ANOVA or Kruskal-Wallis test, P < 0.05). # Statistically significant difference between regions of the gut (ANOVA or Kruskal-Wallis test, P < 0.05).

	[Ti] µmol g ⁻¹ dry mass						
			Oesophagus	Stomach	Mid intestine	Hind intestine	
	Control	Stripped	< 0.003 ^a	< 0.003 ^a	0.004 ± 0.001^{a}	0.005 ± 0.002^{a}	
		Mucosa	0.018 ± 0.009^a	0.010 ± 0.005^a	0.012 ± 0.008^a	0.019 ± 0.012^{a}	
	-	% in mucosa	90.0 ± 14.5^{a}	$83.3\pm18.3^{\rm a}$	75.0 ± 16.9^{a}	79.2 ± 16.8^{a}	
	BulkTiO ₂	Stripped	$0.018 \pm 0.003^{a,*}$	0.030 ± 0.009^{a}	$0.135 \pm 0.039^{ab,\ast}$	$0.252\pm0.131^{\text{b}}$	
Treatments		Mucosa	$0.128 \pm 0.039^{a,*}$	$0.065 \pm 0.019^{a, \ast}$	$0.159 \pm 0.049^{a_{*}}$	$0.148 \pm 0.035^{a,\ast}$	
		% in mucosa	$87.7\pm6.6^{\rm a}$	68.4 ± 5.9^{a}	54.1 ± 3.1^{b}	$37.0\pm10.8^{\text{b}}$	
	TiO ₂ NP	Stripped	$0.019 \pm 0.005^{a, \ast}$	$0.021 \pm 0.004^{ac,*}$	$0.056 \pm 0.010^{b,\ast}$	$0.040 \pm 0.011^{bc,\ast}$	
		Mucosa	$0.174 \pm 0.019^{a,}{*}$	$0.060 \pm 0.006^{b,*}$	$0.136 \pm 0.025^{a,}{*}$	$0.165 \pm 0.026^{a_{*}}$	
		% in mucosa	90.2 ± 2.1^{ac}	74.1 ± 4.3^{bc}	70.8 ± 4.1^{b}	$80.1\pm2.9^{\rm c}$	

Table 3.2 Total Ti metal concentrations in stripped portions of gut and the corresponding mucosa following exposure of isolated whole gut sacs to 1 mg l^{-1} TiO₂ in the gut lumen for 4 h.

Values are means \pm S.E.M. (n = 6 for each group) expressed as total Ti metal concentration in µmol Ti g⁻¹ dry mass of intestinal tissue. Whole gut sacs were filled with physiological saline (control), or the saline containing 1 mg l⁻¹ Bulk TiO₂ or 1 mg l⁻¹ TiO₂ NP at laboratory temperature (18°C) for 4h. The mucosa was stripped off each section of gut so that the mucosa and underlying sub-mucosa/muscularis (stripped gut) separately analysed for total Ti metal concentration. % Ti in the mucosa is calculated for each region of the gut, % Ti in mucosa= [Ti] mucosa/([Ti] stripped + [Ti] mucosa) x 100. * Statistically significant difference from control values within columns (ANOVA or Kruskal-Wallis test, P < 0.05). Different letters (within rows) indicate a statistically significant difference between regions of the gut (ANOVA or Kruskal-Wallis test, P < 0.05). There was no material-type effect (bulk versus nano treatment) within each region of the gut for the tissue Ti concentrations.

[Metal] µmol g ⁻¹ dry mass							
Parameters			Oesophagus	Stomach	Mid intestine	Hind intestine	
Potassium	Control	Stripped	154.03 ± 13.02^{a}	219.29 ± 16.43^{ab}	247.64 ± 43.49^{b}	201.01 ± 35.76^{ab}	
		Mucosa	$201.58 \pm 52.72^{\mathrm{a}}$	164.73 ± 48.37^{a}	$092.72 \pm 22.09^{\mathrm{a}}$	$159.80 \pm 7.90^{\mathrm{a}}$	
		% in mucosa	$52.5\pm6.7^{\rm a}$	39.2 ± 5.4^{ab}	$27.6 \pm 5^{\mathrm{b}}$	$46.5\pm4.5^{\rm a}$	
	Bulk TiO ₂	Stripped	141.55 ± 12.91^{a}	209.97 ± 24.23^{ab}	276.31 ± 16.83^{b}	308.14 ± 63.70^{b}	
		Mucosa	267.92 ± 83.12^{a}	172.09 ± 34.56^{a}	151.30 ± 20.96^{a}	206.89 ± 30.20^a	
		% in mucosa	$59.8\pm6.8^{\mathrm{a}}$	44.5 ± 4.2^{ab}	34.8 ± 2.9^{b}	$42.5\pm6.6^{\rm b}$	
	TiO ₂ NP	Stripped	155.76 ± 15.31^{a}	204.54 ± 8.48^{b}	$230.74 \pm 16.40^{\circ}$	197.41 ± 12.36^{bc}	
		Mucosa	241.83 ± 31.36^{ac}	$118.11 \pm 15.65^{\mathrm{b}}$	$212.13 \pm 41.20^{ab,*}$	$326.88 \pm 40.27^{c,*^{\#}}$	
		% in mucosa	$59.9\pm4.7^{\rm a}$	35.9 ± 2.6^{b}	46.2 ± 3.5^{b}	61.3 ± 3.9^{a}	
Sodium	Control	Stripped	253.18 ± 39.47^{a}	245.68 ± 16.23^{a}	220.82 ± 30.35^{a}	217.07 ± 20.15^{a}	
		Mucosa	424.30 ± 79.28^{ab}	461.25 ± 149.63^{a}	$182.22 \pm 31.90^{\rm b}$	217.07 ± 46.28^{ab}	
		% in mucosa	$60.3\pm6.2^{\rm a}$	$59.4\pm5.8^{\rm a}$	44.9 ± 6.8^{a}	$47.9\pm5.7^{\rm a}$	
	Bulk TiO ₂	Stripped	222.97 ± 019.95^{a}	240.64 ± 027.00^{ab}	333.00 ± 76.18^{ab}	$343.65 \pm 47.26^{b,*}$	
		Mucosa	542.83 ± 119.89^{a}	692.06 ± 177.26^{a}	380.72 ± 117.03^{b}	400.78 ± 147.91^{b}	
		% in mucosa	$67.5\pm5.1^{\rm a}$	71.6 ± 4.2^{a}	51.2 ± 2.9^{b}	$48.4\pm7.5^{\rm b}$	
	TiO ₂ NP	Stripped	213.63 ± 13.01^{a}	$215.54 \pm 05.20^{a,\#}$	$224.85 \pm 15.76^{\rm a}$	$231.61 \pm 36.90^{a,\#}$	
		Mucosa	$516.71 \pm 85.86^{\rm a}$	$422.99 \pm 56.36^{\rm a}$	$444.52\pm 44.42^{a,*^{\#}}$	$528.76 \pm 171.66^{a, \#}$	
		% in mucosa	$69\pm3.4^{\mathrm{a}}$	64.9 ± 3.2^{a}	$65.8\pm2.4^{\mathrm{a}}$	66.2 ± 3.4^{a}	
Calcium	Control	Stripped	$17.20\pm2.89^{\rm a}$	$15.97 \pm 1.91^{\mathrm{a}}$	21.02 ± 4.35^{a}	$21.88\pm7.85^{\rm a}$	
		Mucosa	$24.69\pm5.42^{\rm a}$	$22.74 \pm 6.06^{\mathrm{a}}$	$9.57 \pm 1.28^{\mathrm{b}}$	$12.71 \pm 2.60^{\mathrm{b}}$	
		% in mucosa	$56.3\pm6.2^{\rm a}$	$55.4\pm4.3^{\rm a}$	34.2 ± 5.6^{b}	41.1 ± 6.7^{ab}	
	Bulk TiO ₂	Stripped	$33.09 \pm 18.85^{\mathrm{a}}$	$15.59 \pm 1.19^{\mathrm{b}}$	27.02 ± 3.92^{a}	$29.55 \pm 7.41^{ m a}$	
		Mucosa	$33.23\pm7.72^{\rm a}$	33.16 ± 7.07^{a}	18.30 ± 4.11^{b}	16.32 ± 4.28^{b}	
		% in mucosa	$58.6\pm5.7^{\rm a}$	66 ± 3.7^{a}	39.6 ± 2.9^{b}	38.5 ± 9.7^{b}	
	TiO ₂ NP	Stripped	$22.99\pm9.76^{\rm a}$	13.68 ± 1.61^{a}	17.95 ± 1.76^{a}	15.26 ± 2.18^{a}	
		Mucosa	$28.85\pm3.92^{\rm a}$	$23.48 \pm 3.75^{\mathrm{a}}$	$18.60 \pm 2.02^{a,*}$	23.57 ± 6.30^{a}	
		% in mucosa	60.7 ± 5.6^{a}	61.8 ± 4.6^{ab}	50.7 ± 2.9^{b}	57.9 ± 4.7^{ab}	

Table 3.3 Total K^+ , $Na^+ Ca^{2+}$ and Mg^{2+} concentrations in stripped portions of gut and the corresponding mucosa following exposure of isolated whole gut sacs to 1 mg l⁻¹ TiO₂ in the gut lumen for 4 h.

Table 3.3 continued

			[Metal] µmol g ⁻¹ dry mass			
Parameters			Oesophagus	Stomach	Mid intestine	Hind intestine
Magnesium	Magnesium Control Stripped		$14.74\pm0.96^{\rm a}$	18.08 ± 1.52^{a}	18.54 ± 2.81^{a}	16.66 ± 2.55^{a}
		Mucosa	29.43 ± 7.62^{a}	33.03 ± 11.13^{a}	$11.93\pm2.71^{\text{b}}$	$16.35\pm1.88^{\text{b}}$
		% in mucosa	$61.9\pm5.8^{\rm a}$	$58.6\pm6.2^{\rm a}$	$38.7\pm6.5^{\mathrm{b}}$	50.2 ± 3.9^{ab}
	Bulk TiO ₂	Stripped	$4.52\pm0.92^{\rm a}$	16.61 ± 1.51^{a}	20.98 ± 1.77^{a}	24.27 ± 4.85^{b}
		Mucosa	40.71 ± 13.70^{a}	37.34 ± 10.32^{a}	19.40 ± 3.08^{b}	19.46 ± 2.74^{b}
		% in mucosa	67.4 ± 6.2^{a}	$65.6\pm4.9^{\rm a}$	47.2 ± 4.3^{b}	$46.4\pm6.8^{\rm b}$
	TiO ₂ NP	Stripped	15.60 ± 1.31^{a}	$17.26\pm0.76^{\rm a}$	$17.08\pm1.25^{\rm a}$	$15.54\pm0.90^{\rm a}$
		Mucosa	32.74 ± 3.31^{a}	$23.95\pm2.82^{\mathrm{a}}$	$27.71 \pm 3.92^{a,*^{\#}}$	$31.79 \pm 3.79^{a,*^{\#}}$
		% in mucosa	$67.2\pm3.2^{\rm a}$	57.2 ± 2.8^{b}	61 ± 2.5^{ab}	66.2 ± 3.1^{a}

Values are means \pm S.E.M. (n = 6 for each group) expressed as total electrolytes concentration in µmol g⁻¹ dry mass of intestinal tissue. Whole gut sacs were filled with physiological saline (control), or the saline containing 1 mg l⁻¹TiO₂ NP or 1 mg l⁻¹Bulk TiO₂ at laboratory temperature (18 °C) for 4h. The mucosa was stripped off each section of gut so that the mucosa and underlying sub-mucosa/muscularis (stripped gut) separately analysed for total electrolytes concentration. % electrolytes in the mucosa is calculated for each region of the gut, % electrolytes in mucosa= [electrolytes] mucosa / ([electrolytes] stripped + [electrolytes] mucosa) x 100. * Statistically significant difference from control values or # Statistically significant difference from Bulk TiO₂ value (ANOVA or Kruskal-Wallis test, P < 0.05) (within columns). Different letters (within rows) indicate a statistically significant difference between regions of the gut.

3.3.4 The effect of TiO_2 on the viability of the perfused intestine

All the viability criteria of the perfused intestines were good in general and met in all treatments. Measured by the morphological examination of the intestine (light microscopy of fixed wax sections), LDH activity for both the perfusate and mucosal solution (bath samples), as well as the mucosal solution $[K^+]$, $[Na^+]$, and pH.

3.3.4.1 Histopathology

The histological examination of the perfusate intestine after 4 h showed normal gut morphology (Fig. 3.3). The addition of $1 \text{mg } \text{I}^{-1} \text{TiO}_2 \text{ NP}$ or bulk to the mucosal solution had only minor effects on the epithelium regardless of the gas mixture and only observed in one or two out of six animals in total, with no clear treatment-related effect. These included some slight swelling of the occasional goblet cells (in one out of six animals per treatment), minor lifting of the epithelial cell (in two out of six animals per treatment) and appearance of vacuoles (in two out of six animals per treatment).

3.3.4.2 LDH measurements

The LDH activity in all experiments (n = 6 per treatment; n = 7 for the control and TiO₂ NP treatment gassed with the standard gas mix) in both perfusate and mucosal solutions remained low and within the viability criteria for the preparation (< 1 IU ml⁻¹). The maximum individual perfusate LDH concentrations reached for control, bulk TiO₂ and TiO₂ NP gassed with standard mix gas were 0.71, 0.73 and 0.79 IU ml⁻¹ respectively. For the low CO₂ experiments, LDH activity also remained low with the maximum individual LDH activity reached in the perfusates being 0.76, 0.78 and 0.79 IU ml⁻¹ for control, bulk TiO₂ and TiO₂ NP respectively. However, statistically significant lower values of the cumulative LDH activity for TiO₂ NP (*t*-test, P = 0.001, Fig. 3.4 A and B) were observed compared to the standard gas mix group, and the same situation was seen with the bulk TiO₂ treatment (*t*-test, P = 0.006). For the mucosal LDH, the maximum

individual values reached for the control, bulk TiO2 and TiO2 NP gassed with standard gas mix were 0.54, 0.65 and 0.47 IU ml⁻¹ respectively; while it was 0.35, 0.42 and 0.35 IU ml ⁻¹ gassed with the low CO₂. Bulk TiO₂ showed statistically significant lower mucosal LDH values than the bulk TiO₂ with standard gas mix (*t*-test, P = 0.026, Fig. 3.5 A and B). The cumulative LDH activity for both perfusate and mucosal solutions showed a linear and steady increase over time for all gas mixtures. In the perfusate for the standard gas mixture, the bulk TiO₂ treatment showed a statistically lower cumulative LDH activity by the end of the experiment compared to either control or TiO_2 NP treatments (Kruskal-Wallis test, P = 0.001, Fig. 3.4 A). The same situation was seen with the low CO₂ treatments (Kruskal-Wallis test, P = 0.003, Fig. 3.4 B). There were no statistically significant differences between treatments in the cumulative LDH in the mucosal solutions with either gas mixture (ANOVA, P > 0.05, Fig. 3.5 A and B), except for the bulk TiO₂ gassed with standard gas mix at 240 min, a statistically significant increase in the cumulative LDH was detected in compare to the control and TiO₂ NP (Kruskal-Wallis test, P = 0.032, Fig. 3.5 A). Additionally, a statistically lower of the cumulative LDH was noticed for the TiO₂ NP at 120 min gassed with the standard gas mix compared to the control and bulk TiO_2 (Kruskal-Wallis test, P = 0.02, Fig. 3.5 A).

3.3.4.3 Effects of TiO₂ on the mucosal solution $[K^+]$, $[Na^+]$, and pH

There were no TiO₂-treatment dependent effects on $[K^+]$, $[Na^+]$, or pH of the mucosal solution for most experiments (Table 3.4 and 3.5), which indicate that the intestine was normal with no measurable leak of electrolytes. The measured levels of $[K^+]$, $[Na^+]$ and pH in the fresh physiological saline used in the experiments were 5.86, 146 mmol l⁻¹ and 7.4, respectively, and in comparison there were only micromolar increases in electrolytes above this background concentration in all experiments. For example, the average increase (grand means of both control and treatments) in the $[K^+]$, $[Na^+]$ in the

mucosal solution gassed with low CO₂ were 0.14 \pm 0.03 and 0.79 \pm 0.19 mmol 1⁻¹ respectively (mean \pm S.E.M., n = 12 experiments) over 4 h, while the increase in the mucosal solution gassed with the standard gas mix over 4 h were 0.27 ± 0.04 and 4.49 \pm 0.41 mmol l⁻¹ respectively (mean \pm S.E.M., n = 14 experiments). The grand mean change in pH for the standard gas mixture for all experiments was 0.39 ± 0.03 , indicating the expected normal base secretion of the intestine. However, in the standard gas mix experiments there was slightly more (statistically significant, ANOVA, P =0.03, Table 3.4) base secretion in the nano treatment compared to bulk or controls (mean \pm S.E.M., n = 20 experiments; increases of pH; 0.29 ± 0.07 , 0.43 ± 0.02 and 0.48 \pm 0.04 for control, bulk TiO₂ and TiO₂ NPs, respectively). There were no treatmentdependent changes in net acid excretion into the mucosal solution for the 0.5% CO₂ treatment (Table 3.5), with a grand mean changes of 0.81 ± 0.02 pH units for all treatments combined (mean \pm S.E.M., n = 18 experiments). However, compared to the standard gas mixture, the 0.5% CO₂ had larger increases in pH; 0.83 ± 0.05 , 0.82 ± 0.03 and 0.79 \pm 0.03 for control, bulk TiO₂ and TiO₂ NPs, respectively (mean \pm S.E.M., n =18 experiments) indicated that the pH was more stable with standard gas mix experiment than that found with low CO_2 one.



Figure 3.3 Histology of the perfusate intestine after 4 h perfusion. (a) Control gassed with standard gas mix showing normal intestine with goblet cells (G), (b) 1 mg l⁻¹ bulk TiO₂ gassed with standard gas mix showing appear of vacuoles (V) and lifting epithelium (L), (c) 1 mg l⁻¹ TiO₂ NP gassed with standard gas mix showing lifting of epithelial cell (L), (d) Control gassed with low CO₂ showing normal tissue, (e) 1 mg l⁻¹ bulk TiO₂ gassed with low CO₂ showing minor lifting epithelium (L), and swelling of the goblet cells (G), (f) 1 mg l⁻¹ TiO₂ NP gassed with low CO₂ showing appear of vacuation (V) and lifting of epithelial cell (L). Scale bar = 50 µm, sections were 8 µm thickness and stained with haematoxylin and eosin (n = 6 per treatment; n = 7 for the control and TiO₂ NP treatment gassed with the standard gas mix).





Figure 3.4 The effect of 1 mg 1^{-1} TiO₂ exposure on the cumulative perfusate LDH activity (cumulative sum of international units of LDH activity per minute) over 4 h in perfused intestine at 18°C, (A) control, bulk TiO₂ and TiO₂ NP gassed with standard gas mix, (B) Control, bulk TiO₂ and TiO₂ NP gassed with low CO₂. Values are means ± S.E.M (n = 6 per treatment; n = 7 for the control and TiO₂ NP treatment gassed with the standard gas mix). *Statistically significant difference from the control value (ANOVA or Kruskal-Wallis test, P < 0.05). # Statistically significant difference from bulk TiO₂ value (ANOVA or Kruskal-Wallis test, P < 0.05).





Figure 3.5 The effect of 1 mg l⁻¹ TiO₂ exposure on the cumulative mucosal solution (bath) LDH activity (cumulative sum of international units of LDH activity per minute) over 4 h in perfused intestine at 18°C, (A) control, bulk TiO₂ and TiO₂ NP gassed with standard gas mix, (B) Control, bulk TiO₂ and TiO₂ NP gassed with low CO₂. Values are means \pm S.E.M (n = 6 per treatment; n = 7 for the control and TiO₂ NP treatment gassed with the standard gas mix). * Statistically significant difference from control values (ANOVA or Kruskal-Wallis test, P < 0.05) within each time point. # Statistically significant difference from bulk TiO₂ value (ANOVA or Kruskal-Wallis test, P < 0.05) within each time point.

	Control				TiO ₂ NP		
Time (min)	pH	Bath Na^+ (mmol l^{-1})	Bath K^+ (mmol l^{-1})	рН	Bath Na ⁺ (mmol l^{-1})	Bath K^+ (mmol l^{-1})	рН
20	76 006	140.0 + 0.50	5.0 + 0.02	7.8 + 0.02*	147.4 + 0.6	5 80 + 0.02	7.8 + 0.02*
50	7.0 ± 0.00	148.8 ± 0.38	5.9 ± 0.05	$7.8 \pm 0.03^{+1}$	$14/.4 \pm 0.0$	3.89 ± 0.03	$7.8 \pm 0.02^{+1}$
60	7.6 ± 0.06	149.2 ± 0.66	6.1 ± 0.05	$7.8 \pm 0.04*$	148.8 ± 0.97	5.97 ± 0.03	$7.8\pm0.02*$
90	7.7 ± 0.07	150.8 ± 1.02	6.2 ± 0.05	7.8 ± 0.04	149.8 ± 0.37	$6.02\pm0.02*$	7.8 ± 0.02
120	7.7 ± 0.07	149.6 ± 0.40	6.1 ± 0.07	7.9 ± 0.04	149.8 ± 0.85	6.05 ± 0.06	7.8 ± 0.02
150	7.7 ± 0.08	153.4 ± 2.46	6.3 ± 0.13	$7.9 \pm 0.05*$	150.3 ± 0.48	6.08 ± 0.04	7.8 ± 0.02
180	7.7 ± 0.08	151.4 ± 0.75	6.3 ± 0.11	7.9 ± 0.05	153.3 ± 2.63	6.22 ± 0.16	7.9 ± 0.02
210	7.7 ± 0.08	151.2 ± 0.73	6.3 ± 0.09	$7.9\pm0.05^{\ast}$	150.8 ± 0.95	6.13 ± 0.07	7.9 ± 0.02
240	7.8 ± 0.08	152.2 ± 0.66	6.4 ± 0.11	7.9 ± 0.04	151.0 ± 0.71	6.17 ± 0.08	7.9 ± 0.02

Table 3.4 Total pH, Na^+ and K^+ concentrations in the mucosal solution after exposure of the isolated perfused intestine to 1 mg l^{-1} TiO₂ gassed with 95% O₂: 5% CO₂ for up to 4h.

Values are means \pm S.E.M. (n = 6 for the bulk TiO₂; n = 7 for the control and TiO₂ NP treatment) expressed as mmol l⁻¹ for both Na⁺ and K⁺ in the mucosal solution (bath solution). * Statistically significant difference from control values (Kruskal-Wallis test or *t*-test, P < 0.05) (within rows).

	Control				TiO ₂ NP			
Time (min)	рН	Bath Na ⁺ (mmol l ⁻¹)	Bath K ⁺ (mmol l ⁻¹)	pH	Bath Na^+ (mmol l^{-1})	Bath K ⁺ (mmol l ⁻¹)	pH	
30	7.92 ± 0.08	145.7 ± 0.88	5.87 ± 0.04	7.92 ± 0.08	146.7 ± 0.33	5.92 ± 0.02	7.98 ± 0.07	
60	8.07 ± 0.08	146.0 ± 0.58	5.90 ± 0.02	8.05 ± 0.07	145.7 ± 0.33	5.89 ± 0.01	8.10 ± 0.06	
90	8.17 ± 0.06	146.3 ± 0.33	5.92 ± 0.03	8.13 ± 0.03	146.7 ± 0.33	5.93 ± 0.02	8.17 ± 0.02	
120	8.23 ± 0.07	146.3 ± 0.33	5.94 ± 0.03	8.20 ± 0.04	146.3 ± 0.33	5.92 ± 0.03	8.21 ± 0.04	
150	8.30 ± 0.04	147.3 ± 0.33	5.97 ± 0.02	8.25 ± 0.02	146.3 ± 0.33	5.93 ± 0.02	8.25 ± 0.02	
180	8.35 ± 0.04	148.0 ± 0.00	6.10 ± 0.06	8.30 ± 0.03	147.3 ± 0.33	6.05 ± 0.08	8.33 ± 0.02	
210	8.38 ± 0.05	147.7 ± 0.33	6.17 ± 0.17	8.33 ± 0.02	147.3 ± 0.33	6.05 ± 0.08	8.33 ± 0.02	
240	8.42 ± 0.04	148.0 ± 0.00	6.23 ± 0.19	8.36 ± 0.02	147.0 ± 1.00	6.14 ± 0.18	8.38 ± 0.02	

Table 3.5 Total pH, Na^+ and K^+ concentrations in the mucosal solution after exposure of the isolated perfused intestine to 1 mg l^{-1} TiO₂ gassed with 99.5% O₂: 0.5% CO₂ for up to 4h.

Values are means \pm S.E.M. (n = 6 for each group) expressed as mmol l⁻¹ for both Na⁺ and K⁺ in the mucosal solution (bath solution). There was no statistically significant difference (Kruskal-Wallis test or *t*-test, P > 0.05) between groups (control, bulk or nano) within each time point over 4 h perfusion.

Mucosal TiO ₂ (mg Γ^1)								
		Gassed with 95% O ₂ : 5%	CO_2		Gassed with 99.5% O ₂ : 0.5% CO ₂			
Time (min)	Control	Bulk TiO ₂	TiO ₂ NP	Control	Bulk TiO ₂	TiO ₂ NP		
30	< 0.004	$0.37\pm0.05*$	$0.10\pm0.02^{\#}$	< 0.004	$0.16\pm0.01*$	$0.18\pm0.07*$		
60	< 0.004	$0.32 \pm 0.03*$	$0.19\pm0.07*$	< 0.004	$0.08\pm0.03^{\ast}$	$0.21 \pm 0.13*$		
90	< 0.004	$0.36\pm0.02*$	$0.12 \pm 0.04^{*^{\#}}$	< 0.004	$0.09 \pm 0.03*$	$0.21\pm0.12*$		
120	< 0.004	$0.34\pm0.04*$	$0.17 \pm 0.05 *^{\#}$	< 0.004	$0.08\pm0.03^*$	$0.13\pm0.08*$		
150	< 0.004	$0.35\pm0.03*$	$0.18 \pm 0.10^{\ast \#}$	< 0.004	$0.05\pm0.02*$	$0.05\pm0.03^{\ast}$		
180	< 0.004	$0.34 \pm 0.04*$	$0.09 \pm 0.02^{*^{\#}}$	< 0.004	$0.03\pm0.01^*$	$0.07\pm0.03^{\ast}$		
210	< 0.004	$0.35\pm0.02*$	$0.14 \pm 0.04^{*^{\#}}$	< 0.004	$0.03\pm0.01^*$	$0.07\pm0.03^{\ast}$		
240	< 0.004	$0.38\pm0.01*$	$0.16 \pm 0.05 *^{\#}$	< 0.004	0.02 ± 0.01	$0.06 \pm 0.03^{*}$		

Table 3.6 Total TiO_2 concentration in the mucosal solution after exposure of the isolated perfused intestine to 1 mg l⁻¹ TiO_2 for up to 4h.

Values are means \pm S.E.M. (n = 6 per treatment; n = 7 for the control and TiO₂ NP treatment gassed with the standard gas mix) expressed as mg l⁻¹ for TiO₂ in the mucosal solution (bath solution) gassed with 95% O₂: 5% CO₂ and 99.5% O₂: 0.5% CO₂. * Statistically significant difference from control values or # Statistically significant difference from Bulk TiO₂ value (ANOVA or Kruskal-Wallis test, P < 0.05) (within columns).

3.3.5 TiO₂ Distribution in the mucosal solution

The dispersion of TiO₂ was confirmed by the nanoparticles tracking analysis system (Nanosight LM 10) which analysed the mucosal solution samples that taken at the beginning (at 30 min) and at the end (at 240 min) of the experiments. The bulk TiO₂ material remained stable over the 4 h perfusions, at the start of experiments the bulk TiO₂ had a mean particle size of 301.50 ± 43.47 nm, with the smallest size bin in the distributions being 70.61 \pm 14.79 nm (mean \pm S.E.M., n = 6, Fig. 3.6. C). The mean values for the bulk TiO₂ at the end of the experiments were 236.67 ± 24.05 nm, with the smallest size bin of 61.5 ± 7.10 nm (mean \pm S.E.M., n = 6, neither being statistically different from the start of the experiment, t-tests, P > 0.05, Fig. 3.6. D). TiO₂ NP showed statistically significant lower mean particle size values (t-tests, P < 0.05) compared to the bulk TiO_2 treatment at each time point. The mean size of the TiO_2 NP at the beginning of the experiment and after 4 h perfusion were 169.43 ± 18.77 nm and 140.18 ± 15.85 nm (mean \pm S.E.M., n = 10, t-test, P = 0.84), respectively, while the smallest size bin of particles had a mean of 50.10 ± 3.71 nm and 46.93 ± 1.88 nm (mean \pm S.E.M., n = 10, t-test, P = 0.66), at the start and end of the experiment respectively (see Fig. 3.6. A and B).



Figure 3.6 The dispersion of 1 mg 1^{-1} TiO₂ in the mucosal solution. Nanosight graphs are represent examples of individual samples from replicated experiments (TiO₂ NP, *n* = 10 and Bulk TiO₂, *n* = 6). (A) TiO₂ NP distribution in mucosal solution at the beginning of the experiment (at 30 min), mean are 103 nm, started with 26 nm primary particle size. (B) TiO₂ NP distribution in mucosal solution at the end of the experiment (at 240 min), mean are 81 nm, with 28 nm primary particle size. (C) Bulk TiO₂ in mucosal solution at the beginning of the experiment (at 30 min), mean are 136 nm and 36 nm primary particle size. (D) Bulk TiO₂ in mucosal solution at the end of the experiment (at 240 min), mean are 122 nm with 65 nm primary particle size.

3.3.6 Ti uptake from TiO_2 by the perfused intestine

3.3.6.1 Titanium accumulation from TiO_2 exposures in the perfused intestine

The accumulation of Ti from TiO_2 in the tissue during the perfusions is shown in Fig. 3.7. As expected, total Ti metal concentrations in the tissues were higher in both of the TiO₂ treatments compared to the no added TiO₂ control. However, there were differences between the mid and hind gut depending on both types of TiO₂ treatment, and the gas supplement used. In general, there was a material-type effect, with TiO_2 NP treatment showing lower total Ti metal concentrations in the tissue than the equivalent bulk treatment when using the standard gas mixture (Fig. 3.7. A). The hind gut also generally accumulated less Ti than the mid gut within treatment (statistically significant, *t*-tests, P < 0.05, Fig. 3.7. A). For the mid gut gassed with standard gas mix the total Ti metal concentrations in control, bulk TiO₂ and TiO₂ NP groups were $< 0.004, 0.094 \pm$ 0.041 and 0.019 \pm 0.004 µmol g⁻¹ dry weight of tissue, respectively (mean \pm S.E.M., n =6 perfusions for the bulk TiO₂; n = 7 perfusions for the control and TiO₂ NP treatment, Kruskal-Wallis test, P = 0.002). In the hind gut the total Ti metal concentrations in control, bulk TiO₂ and TiO₂ NP groups were 0.003 \pm 0.001, 0.022 \pm 0.005 and 0.009 \pm 0.003 μ mol g⁻¹ dry weight of tissue, respectively (mean \pm S.E.M., n = 6 perfusions for the bulk TiO₂; n = 7 perfusions for the control and TiO₂ NP treatment, Kruskal-Wallis test, P = 0.02). Bulk TiO₂ showed significantly higher values than that found in both control and TiO₂ NP groups for either mid or hind gut.

The addition of low CO₂ into the mucosal solution changed the Ti accumulation in the tissue compared to the standard gas mix groups (Fig. 3.7. B); showing higher Ti accumulation with TiO₂ NP treatment than the equivalent bulk treatment. The Ti concentration from TiO₂ in the mid gut for control, bulk TiO₂ and TiO₂ NP groups were 0.002 ± 0.001 , 0.013 ± 0.005 and $0.029 \pm 0.009 \ \mu mol g^{-1}$ dry weight of tissue, respectively (mean \pm S.E.M., n = 6, Kruskal-Wallis test, P = 0.003). The hind gut Ti concentrations were 0.002 ± 0.001 , 0.009 ± 0.002 and $0.023 \pm 0.009 \mu mol g^{-1}$ dry weight of tissue in control, bulk TiO₂ and TiO₂ NP groups respectively (mean ± S.E.M., n = 6, Kruskal-Wallis test, P = 0.003). The TiO₂ NP treatment was significantly higher than the bulk TiO₂ one. No significantly differences were found for the Ti concentration between the mid and hind intestine (*t*-test, P > 0.05) in all treatment groups.



Figure 3.7 Total Ti metal concentrations in the tissues determined by ICP-OES in the mid (blue bars) and hind gut (red bars) after 4 h perfusions with a concentration of 1 mg Γ^1 TiO₂ (bulk or NP) added to the mucosal solution gassed with standard gas mix (A) or low CO₂ (B). Data are means ± S.E.M. (n = 6 perfusions per treatment; n = 7 perfusions for the control and TiO₂ NP treatment gassed with the standard gas mix). * Statistically significant difference from control values (ANOVA or Kruskal-Wallis test, P < 0.05). # Statistically significant difference from bulk TiO₂ treatment (ANOVA or Kruskal-Wallis test, P < 0.05). Letter (a) indicates statistically significant differences between mid and hind intestine (*t*-test, P < 0.05).

3.3.6.2 Net Ti flux to the serosal compartment

The cumulative Ti metal uptake across the intestine into the serosal perfusate for individual experiments at different size (NP and Bulk) and different gas supplement (standard gas mix and low CO₂) are shown in Fig. 3.8. There was a steady appearance of Ti in the eluted perfusate over time in the treated animals compared to controls (Fig. 3.8). There was also a substantial difference between the $TiO_2 NP$ accumulations in the perfusate associated with the type of gas mixture. Changing from the standard gas mixture to the low CO₂ gas mixture greatly enhanced the appearance of Ti in the serosal perfusate from both the bulk and nano TiO₂ exposures (compare bulk, Fig. 3.8.B and E; compare nano, Fig. 3.8.C and F). The maximum initial uptake rate of NP into the serosal perfusate with standard gas mix showed a lower values by 13 folds than the low CO₂ being 1.55 ± 0.33 and 21.16 ± 18.90 nmol g⁻¹ h⁻¹ (mean \pm S.E.M., n = 6 perfusions per treatment; n = 7 perfusions for the control and TiO₂ NP treatment gassed with the standard gas mix), respectively. The bulk TiO₂ showed low content of Ti in the serosal perfusate being 0.98 ± 0.47 nmol g⁻¹ h⁻¹ for the standard gas mix (mean \pm S.E.M., n = 6) and 3.42 ± 0.74 nmol g⁻¹ h⁻¹ for the low CO₂ (mean ± S.E.M., n = 6). Ti content in the serosal perfusate from bulk TiO₂ represented a significantly lower values than Ti from TiO₂ NP group for both standard gas mix (t-test, P = 0.04) and low CO₂ (t-test, P =0.001). However, reduce the appearance of the Ti in the serosal perfusate did not compromise the steady perfusate flow (one of the viability criterion for the preparation).



Figure 3.8 The cumulative appearance of total Ti metal in the serosal compartment (red bars, nmol Ti metal) of perfused intestine preparations from rainbow trout exposed to 1 mg Γ^1 of TiO₂ in the mucosal saline. Cumulative perfusate flow (blue diamonds, ml) is also shown. Graphs are representative examples of individual perfusions from replicated experiments (n = 6 perfusions per treatment; n = 7 perfusions for the control and TiO₂ NP treatment gassed with the standard gas mix). (A) Control with no added TiO₂, (B) bulk TiO₂, (C) TiO₂ NPs, (D) control gassed with low CO₂, (E) bulk TiO₂ gassed with low CO₂; (F) TiO₂ NPs gassed with low CO₂. Fish weights in the examples are 307, 136.3, 249.9, 229.8, 648.9 and 140.3g, respectively.

3.3.6.3 Perfusate Ti from TiO₂ and water uptake rates

Calculated net Ti from TiO₂ uptake rates and water flux across the intestine are shown (Table 3.7). When using the standard gas mixture, both bulk and nano TiO₂ treatments showed a trend of higher net fluxes compared to the no-added TiO₂ controls. However, only the TiO₂ NP treated showed a statistically significant higher net flux to the serosal compartment (7 and 9 fold higher than controls for initial and overall flux rates respectively, Table 3.7). For the standard gas mixture, there were no differences between the initial and overall Ti flux rates, indicating that the flux measurements were not hampered by saturation of the Ti electrochemical gradient (still movement of Ti across the tissue). Importantly, these net fluxes of Ti were in the opposite direction to the net water flux across the tissue (not explained by passive solvent drag). Both the bulk and TiO₂ NP treatments increased the net outward flux of water across the intestine compared to the control, and there was a statistically significant material-type effect with the bulk treatment causing more net water efflux than the nano treatment for both initial and overall rates (Kruskal-Wallis test, *P* < 0.05, Table 3.7).

3.3.7 Tissue electrolytes and moisture content of perfused intestine

Tissue electrolytes concentrations and moisture content of the perfused intestine are shown (Table 3.8). Using the standard gas mixture, there was generally no TiO₂ treatment effect on tissue Na⁺ concentrations, but there was a statistically significant increase (Kruskal-Wallis test, P < 0.05) in tissue K⁺, Mg²⁺ (in both the mid and hind intestine), and Ca²⁺ (just in the hind intestine) for the bulk TiO₂ in comparison with the control; and compared to TiO₂ NP treatments indicating a material-type effect (Table 3.8). The moisture content of the intestine was not affected by any treatment (Table 3.8).

3.3.8 Effects of lowering CO₂ to 0.5%

Some of the largest changes in total Ti metal concentration in the tissues and Ti uptake observed in these experiments were due to changing the gas mixture for the standard 95% O₂: 5% CO₂ to 99.5% O₂: 0.5% CO₂. In general, the effect of using the lower CO₂ mixture was to cause large increases in the net Ti flux to the serosal compartment (Table 3.7, Fig. 3.8), associated with either depletion (bulk material) or elevation (nano material) of the total Ti metal concentration in the tissue compared to the standard gas mixture (Fig. 3.7B). Most notably for the TiO₂ NP treatment, the initial Ti flux rate increased from 1.6 to 21.2 nmol g⁻¹ h⁻¹ (13 fold increase) when the lower CO₂ mixture was used (Table 3.7). This was accompanied by an increase in the total Ti metal concentration in the tissue in the 0.5% CO₂ treatment containing TiO₂ NPs (mean \pm S.E.M., *n* = 6, mid intestine, 0.03 \pm 0.01; hind intestine, 0.02 \pm 0.01 µmol g⁻¹dry weight, Fig. 3.7) equating to about a 1.5 and 2.5 fold increase in these tissue respectively due to the lower CO₂ gas mixture.

There was a strong effect associated with the type of material and the increase in the initial flux rate, due to changing the gas mixture, for the bulk material was from 0.98 to 3.4 nmol g⁻¹ h⁻¹ (Table 3.7). This represents a 3.5 fold increase in net Ti flux, but even this is small compared to the 13 fold increase with the equivalent nanomaterial treatment (statistically significant material-type effect, Table 3.7). For the bulk material, unlike the nano scale TiO₂, the CO₂ mixture effect was to decrease the total Ti metal concentration in the tissue (e.g., a 7.2 fold decrease in the mid intestine, Fig. 3.7B). Changing the gas mixture had no effect on unexposed control intestines, and there were no anatomical regional differences for the gut in terms of the Ti accumulation response to the 0.5% CO₂ gas mixture (*t*-test, P > 0.05, Fig. 3.7B).

Altering the gas mixture had no statistically significant effects on net water fluxes in the intestines from control fishes, or from guts in the bulk TiO_2 treatment.

Unlike the standard gas mix the large rise in net water efflux associated with exposure to the TiO_2 NP material (Table 3.7), indicating a material-type effect of the CO_2 manipulation on water fluxes.

Changing the gas mixture generally had no effect on the electrolytes in the intestines from unexposed control animals, but did cause an increase (not significantly difference) of K^+ , Ca^{2+} and Mg^{2+} concentrations in the mid intestine from the TiO₂ NP treatment, but notably in this treatment, the hind intestine also showed depletion of the same electrolytes (Table 3.8). Changing the gas mixture in the bulk TiO₂ treatment generally caused much larger increases in tissue Na⁺, K⁺ and Ca²⁺ concentrations than the equivalent nano treatment (combined gas and material-type effect on tissue electrolytes, Table 3.8). Moisture content and Mg²⁺ concentrations were generally unaffected by changing the gas mixture (Table 3.8).

3.3.9 Presence of particles within the gut mucosa

Transmission electron microscopy images of the gut epithelium from control, bulk and TiO_2 NP treated fish are shown in Fig. 3.9 and 3.10. For all treatments, except the no added particle control, electron dense particles are observed inside the epithelial cells with some of the particles observed inside vesicles, or associated with tight junctions (Fig. 3.9 and 3.10). Damage of the mitochondria (Fig. 3.10) was shown in the gut tissue after exposure to the bulk TiO_2 represented by degradation with lose of the structure of the mitochondria. The damaged mitochondria after exposure to TiO_2 NP were exhibited by a loss of the structure of the cristae inside the mitochondria. Degeneration of the microvilli was also noted in the epithelial cells after being exposed to both bulk and TiO_2 NP treatments (Fig. 3.9).

Mucosal [TiO ₂]	Net Ti flux	x, $J_{\rm net,Ti}$	(ml g ⁻¹ h ⁻¹) Net water flux, $J_{net,H2O}$		
$(1 mg l^{-1})$ and Drugs	(nmol g	⁻¹ h ⁻¹)			
Concentration	Initial rate	Overall rate	Initial rate	Overall rate	
Control	0.23 ± 0.16	0.26 ± 0.04	-6.22 ± 0.85	-7.50 ± 1.18	
Bulk TiO ₂	0.98 ± 0.47	0.85 ± 0.32	$-26.28 \pm 2.52*$	$-28.54 \pm 3.19*$	
TiO ₂ NP	$1.55 \pm 0.33^{*\#}$	$2.38 \pm 0.68 *^{\#}$	$-17.56 \pm 4.00^{*\#}$	$-18.01 \pm 4.16^{*^{\#}}$	
0.5% CO ₂ Control	0.51 ± 0.14	0.06 ± 0.03	-3.42 ± 1.71	-3.97 ± 0.97	
0.5% CO ₂ Bulk TiO ₂	$3.42\pm0.74^{\ast\ddagger}$	$3.18 \pm 0.42^{*\ddagger}$	$-7.05 \pm 2.29^{\ddagger}$	$-8.95 \pm 1.78^{*\ddagger}$	
0.5% CO ₂ TiO ₂ NP	$21.16 \pm 18.90^{*^{\# \ddagger}}$	$3.27 \pm 1.60^{*a}$	$-17.13 \pm 2.37^{*^{\#}}$	$-22.30 \pm 6.40^{*^{\#}}$	

Table 3.7 Effects of exposure to $1 \text{mg } \Gamma^1 \text{TiO}_2$ with/without different gases on the Ti and water fluxes across the isolated perfused trout intestine.

Values are means \pm S.E.M. (n = 6 perfusions per treatment; n = 7 perfusions for the control and TiO₂ NP treatment gassed with the standard gas mix). expressed per gram dry mass of intestine per hour. Negative values indicate a net loss from the serosal solution, initial and overall rates data were calculated from cumulative perfusate data at 10 min and 4 h, respectively.* Statistically significant difference from the relevant control values within columns (ANOVA or Kruskal-Wallis test, P < 0.05), # Statistically significant difference from the relevant control values within columns (ANOVA or Kruskal-Wallis test, P < 0.05). Letter (a) within rows indicates a statistically significant difference between initial and overall rates (*t*-test, P < 0.05).

Mucosal [TiO ₂]	Tissue [Metal] µmol g ⁻¹ dry mass							
$(1 \text{mg } l^{-1})$	Na ⁺	\mathbf{K}^+	Ca ²⁺	Mg^{2+}	Moisture %			
Mid Intestine								
Control	145.5 ± 11.9	142.9 ± 16.0	17.0 ± 4.5	14.0 ± 1.7	81.0 ± 1.8			
Bulk TiO ₂	156.1 ± 24.3	$212.0 \pm 15.8*$	18.6 ± 3.5	$20.2\pm1.7*$	78.4 ± 2.1			
TiO ₂ NP	162.8 ± 34.6	$136.1 \pm 18.3^{\#}$	12.5 ± 1.3	$12.1 \pm 1.2^{\#}$	82.3 ± 1.7			
0.5% CO ₂ Control	127.8 ± 11.5	145.6 ± 14.2	15.1 ± 3.8	13.0 ± 0.9	80.5 ± 1.2			
0.5% CO ₂ Bulk TiO ₂	$329.1 \pm 26.1*$	$283.3 \pm 13.8*$	$35.2 \pm 3.1*$	16.3 ± 1.7	82.6 ± 0.8			
$0.5\% \text{ CO}_2 \text{TiO}_2 \text{ NP}$	$118.8 \pm 17.9^{\#}$	$161.9 \pm 24.6^{\#}$	$22.9\pm9.7^{\#}$	14.3 ± 2.6	83.4 ± 1.8			
Hind Intestine								
Control	117.4 ± 7.6	127.2 ± 7.9	9.5 ± 0.6	10.8 ± 0.4	81.6 ± 1.4			
Bulk TiO ₂	116.5 ± 7.6	$168.3 \pm 5.9*$	$18.0 \pm 4.2*$	$15.8\pm0.7*$	81.5 ± 1.8			
TiO ₂ NP	98.7 ± 9.5	$121.0 \pm 7.1^{\#}$	$7.9\pm0.5^{\#}$	$9.9\pm0.6^{\#}$	80.9 ± 1.4			
0.5% CO ₂ Control	109.3 ± 8.8	135.1 ± 13.1	9.6 ± 0.9	11.0 ± 0.9	81.1 ± 1.6			
0.5% CO ₂ Bulk TiO ₂	$399.7 \pm 25.8*$	$293.3 \pm 11.1*$	$21.1 \pm 2.4*$	17.9 ± 1.1	84.4 ± 0.6			
$0.5\% \text{ CO}_2 \text{TiO}_2 \text{ NP}$	$75.1\pm8.6^{\#}$	$107.8 \pm 17.8^{\#}$	14.5 ± 5.0	$9.5\pm1.3^{\#}$	78.5 ± 3.5			

Table 3.8 Total K⁺, Na⁺, Ca²⁺, and Mg²⁺ Concentrations of gut tissue following exposure of isolated perfusate trout intestine to 1 mg l⁻¹ TiO₂ for 4 h.

Values are means \pm S.E.M. (n = 6 perfusions per treatment; n = 7 perfusions for the control and TiO₂ NP treatment gassed with the standard gas mix). expressed as μ mol g⁻¹ dry mass of intestinal tissue, except for moisture content (%) = ((wet weight – dry weight)/wet weight) x 100. * Statistically significant difference from control values within columns (ANOVA or Kruskal-Wallis test, P < 0.05), # Statistically significant difference from bulk TiO₂ value within columns (ANOVA or Kruskal-Wallis test, P < 0.05).
Chapter 3



Figure 3.9 TEM images of rainbow trout gut epithelium cells. (a,b) control showing normal structure of the mitochondria (Mt); (c,d) 1 mg I^{-1} bulk TiO₂ showing accumulation of assumed bulk TiO₂ particles (Bk) and some damaged mitochondria (Mt); (e,f) 1 mg I^{-1} TiO₂ NP show examples of assumed TiO₂ nanoparticles (NP) inside the vacuoles and near to the tight junction with some damaged mitochondria (Mt). (c) and (e) Images show some occasional fusion or degeneration of parts of the villi (Dg). Scale bar = 2 µm in (a, c and e) and 1 µm in (b, d and f) as well as 500 nm, (*n* = 3 per treatment). Composition of the particles was not confirmed in the electron micrographs.



Figure 3.10 TEM images of rainbow trout gut epithelium cells (a) control showing normal structure of mitochondria (Mt), (b) 1 mg l⁻¹ bulk TiO₂ showing some damaged mitochondria (Mt) represented by degradation with lose of the structure of the mitochondria, (c) and (d) 1 mg l⁻¹ TiO₂ NP showing some damaged mitochondria (Mt) represented loss of the structure of the cristae inside the mitochondria. Scale bar = 1 μ m in (a, b and d) as well as 500 nm, and 2 μ m in (c), (*n* = 3 per treatment).

3.4 Discussion

This is the first reports on transepithelial uptake of NPs by the gut of rainbow trout, using the isolated gut perfusion technique. TiO_2 is one of the most widely used NPs and this is the first study designed to show how particles cross the gut, after the evidence of Ti accumulation in the gut, liver, gill, brain and spleen of trout (Federici et al., 2007; Ramsden et al., 2009).

3.4.1 Where is the TiO_2 NP absorbed along the gut?

Previous studies have shown Ti accumulation in rainbow trout gut from ingested water containing TiO₂ NPs during aqueous exposure studies (Federici et al., 2007) and from dietary inclusions of TiO₂ NPs (Ramsden et al., 2009). In rodents, Ti accumulation from oral exposure to bulk TiO₂ has also been reported. For example, oral gavage of high mg doses (12.5 mg kg⁻¹) of the rutile TiO₂ form of the bulk material (500 nm size) in rats causes Ti metal accumulation in the internal organs, and notably in the gut associated lymphoid tissue (Jani et al., 1994). However, to our knowledge the regional differences in total Ti metal concentrations in the tissue associated with exposure to TiO₂ particles in the gut have not been previously reported in fish. In this study following treatment with either the bulk or nanomaterial, Ti accumulated in all the regions of the gut from the oesophagus to the hind intestine. For the bulk material treatment, most of the Ti accumulation was in the intestinal regions rather than the stomach (Table 3.1), with the mid intestine having the highest total Ti metal concentrations. However, the situation was different for the TiO_2 NP treatment which showed an even accumulation of Ti in the different parts of the gut, but with a tendency for the total Ti metal concentrations to be higher in the mid and hind gut (not statistically different between gut regions, Table 3.1). There were no major fluid shifts within the gut tissue, with normal water content (66 - 81%) in the different regions of the gut and the absence of disturbances to the major electrolyte levels, especially Na⁺ which is crucial to osmoregulation in the tissue (Table 3.1). This confirms the tissue was not leaking and that the Ti accumulation cannot be explained by osmotic solvent drag. Overall, these data suggest the intestine is more important for Ti uptake from TiO₂ exposures than other parts of the gut, although the form of the accumulated Ti (as particles and/or dissolved Ti) cannot be determined from the tissue concentration measurements alone. Nonetheless, the involvement of the intestine more than other regions of the gut is consistent with reports on many other trace metals (Handy et al., 2000; Hoyle and Handy, 2005). Notably, for both experimental treatments of TiO₂, the measured total Ti metal concentrations were higher in the mucosa, rather than the muscularis (Table 3.2). This suggests a specific accumulation of either Ti metal and/or TiO₂ particles into the epithelial cells (see below). The proportions of the total Ti metal concentration in the mucosa (typically 70% or more) are broadly similar to previous reports for trace metals in fish gut (e.g. Cu in catfish gut, Handy et al., 2000; Hg in trout gut, Hoyle and Handy, 2005).

3.4.2 Viability of the perfused intestine

The viability criteria normally employed for the perfusion techniques (Handy et al., 2000). In the present study, the histology integrity of the gut as a barrier, perfusate and mucosal LDH activity, leak of ions into the mucosal solutions ($[K^+]$, $[Na^+]$) and perfusate flow were used to assess the viability. These parameters are essential to evaluate the viability criteria of the perfusion techniques (Perry et al., 1983; Perry et al., 1984; Handy et al., 2000), and the accepted results should be obtained only from those perfusions which passed all these checks. Current results from this study indicate the suitability of using the perfusion technique with the NPs which is met with the viability criteria of the preparation.

3.4.2.1 Histopathology

Histological examinations of guts after exposure to either bulk or nano TiO_2 showed normal tissue for the all gut regions. However, some minor effects on gut morphology were shown in one or two animals considered by minor lifting of epithelium, swelling of the goblet cells and occasional appearance (occurring at infrequent parts of the tissue) of some vacuoles which would not compromise the intestinal barrier (Fig. 3.3).

3.4.2.2 LDH measurement

One of the most sensitive indicators of cell injury is the LDH activity and the amount of LDH in the normal cells (no damage) compared to that present in the blood stream (Moss and Henderson, 1986; Lott and Nemensanszky, 1987). In trout, the blood normally contains about 1 IU ml⁻¹ LDH *in vivo* (Hille, 1982). In the perfusate, LDH activity is considered as a very sensitive viability indicator (Campbell et al., 1999; Handy et al., 2000), and in this study the LDH activity was far below that found *in vivo* (< 1 IU ml⁻¹) in all experiments, in perfusates (Fig. 3.4) and mucosal solutions (Fig. 3.5). The low LDH leak was evident even when the gas mixture was changed (5% or Low CO₂).

3.4.2.3 [K+], [Na+] and pH content in the mucosal solution

Leak of electrolytes from the tissue was negligible showed only micromolar changes in mucosal solution K^+ , and Na^+ concentration comparing to controls (Table 3.4 and 3.5). The steady rates of eluted perfusate flow suggested good viability of the preparations (Fig. 3.8). Taken together, the good perfusate flow, low LDH leak, and normal electrolytes suggests that the preparations were of good viability and has a utility for measuring the uptake of Ti from both bulk and nano forms of TiO₂.

In the normal medium (5% CO₂), the steady mucosal solution pH and no leakage of the electrolytes also suggests no acidification of the tissue (see Handy et al., 2000; Holye and Handy, 2005). However, decreasing the gas supplement to 0.5% CO₂ caused an elevation of the pH in the mucosal solution from [insert pH] of 7.4 to about pH 7.9, suggesting either the uptake of H^+ ions by the tissue, or the efflux of bicarbonate to result in a more alkaline pH. The secretion of bicarbonate by fish, resulting in decreased serosal HCO_3^- concentration (Grosell et al., 2005), may explain the pH change here. It is interesting that the role of this bicarbonate secretion in fish is to ensure the precipitation of magnesium and calcium carbonates (e.g., MgCO₃), which functions to maintain the outward diffusion gradients for divalent ions (Grosell et al., 2005). The generation of CO₃ may also occur by a spontaneous chemical reaction on the surface of the nanoparticle when the CO₂ interacts with the alkali atoms on TiO₂ (Krischok et al., 2002). This could therefore alter, the HCO₃⁻ equilibrium across the tissue, and therefore explain the effect of lowering the CO₂ in the presence of TiO₂ NPs.

3.4.3 Ti exposure and the distribution of the TiO_2 in the mucosal solution

The exposure to TiO₂ was confirmed by applying different processes to measure the mass concentration of TiO₂ and particle size distributions in the mucosal solution by using nanoparticle tracking analysis (NTA, Fig. 3.6), the appearance of particulate material in the mucosa by electron microscopy (Fig. 3.9 and 3.10), as well as total Ti metal concentration in the intestinal tissue (Fig. 3.7 A and B). NTA measurements were made at 30 and 240 minutes during the experiments using a triplicate samples for each time. In general, the distribution of NP and bulk TiO₂ was good, with the mean particle diameter changing from only 301 to 236 nm for the bulk material, and from 169 to 140 nm for the NPs. The distribution of particles moving towards slightly smaller sizes in the mucosal solution, are probably just a reflection of the faster settling of larger aggregates as proposed by other studies (Keller et al., 2010), and should not be interpreted as selective uptake of larger particles by the tissue. The concentration of TiO_2 in the mucosal solution was 0.35 and 0.14 mg l⁻¹ of bulk and TiO_2 NPs, respectively (Table 3.6) after 4 h perfusions and exposure to 1 mg l^{-1} of TiO₂ suggesting loss of metal and/or particles from the mucosal solution. This phenomena is well known for dissolved metals where the intestinal bioavailability is a fractions of the exposure dose, and is associated with the normal protective role of mucus secreted by the preparation which will chelate then precipitate metal ions (e.g., for Cu perfusions, 77% of the added Cu is lost at exposures of 100 μ mol 1⁻¹, Handy et al., 2000). Obvious clumps of white-coloured mucous precipitate appeared in the bath over time (as observed with gill mucus during waterborne exposures with TiO₂, Federici et al., 2007); suggesting precipitation of particles with the mucus.

3.4.4 The uptake of Ti from TiO_2 by the perfused intestine

3.4.4.1 Tissue TiO₂ uptake

The increase of measured total Ti metal concentrations in the tissue following exposure to either the bulk or nano form of TiO₂ confirmed that the exposure took place, and that Ti (the form cannot be identified from the total tissue Ti measurements) was presents in/on the gut tissue after 4 h perfusion. Most of the metal was internalised with the mid gut generally accumulating more Ti than the hind gut (Fig. 3.7 A and B). Particulate material was also observed inside the epithelial cells using electron microscopy (Fig. 3.9 and 3.10). Similar observations have been made in rat gut associated lymphoid tissue with measurable Ti metal from rutile TiO₂ appearing in/on the tissue (500 nm, dose 12.5 mg kg⁻¹ for 10 days, Jani et al., 1994). Galloway et al. (2010) also recently reported particles from TiO₂ NP exposures in close association with the microvilli in marine polychaete worms.

The intestinal Ti accumulation was greater from the bulk TiO_2 compare to nano exposure in both mid and hind intestine with different gas supplements; suggesting that the bulk TiO_2 was accumulated in the tissue rather than diffuse to the serosal side. However, this does not imply a "nano effect" on tissue Ti accumulation *per se*. This result could be explained by the fact that the bulk material maintained a slightly higher concentration in the mucosal solution than the nano form, or that particle for particle, more Ti on a mass basis is inevitably moved into the tissue by uptake of a bulk particle compared to a smaller nano one. Regardless of any nano effect, the fact that Ti accumulates in the intestine suggests that the efflux of Ti from the gut to the blood must be slower than the mucosal (apical) uptake in order to achieve net accumulation in the tissue.

3.4.4.2 Serosal TiO₂ uptake

In all experiments, there was a cumulative increase of total Ti metal in the serosal perfusate suggesting transepithelial absorption of either Ti metal and/or particulate TiO₂. The nano form of TiO₂ showed higher values compare to bulk TiO₂ in the serosal perfusate content which indicated the transfer of the NP into the serosal perfusate rather than accumulation in the intestinal tissue (i.e., more transepithelial uptake of the nanoform). The steady, and sometimes saturable, appearance of Ti in the eluted perfusate over time (Fig. 3.8), suggests a carrier-mediated mechanism(s). This hypothesis is supported by net water flux being in the opposite direction to net Ti fluxes (Table 3.7) and the absence of dilution of electrolytes in the tissue (Table 3.8); excluding passive solvent drag or leak (see Handy et al., 2000) of any (theoretically) dissolved Ti.

3.4.4.3 Perfusate Ti from TiO₂ and water uptake rates

The differences between the initial and overall TiO_2 uptake rates (Table 3.7) indicate that the serosal solution was at equilibrium with TiO_2 during the experiments. However, this is not simply explained by diffusion process associated with solvent drag of water across the epithelium. First of all this is not a valid scientific argument as NPs are not in solution, but in suspension. There is no logical reason why solvent drag would occur. Also, the water fluxes are in the opposite direction to the Ti fluxes. The latter is best explained by a carrier-mediated process, not by the solvent drag. The same suggestion has been made by Handy et al. (2000) after exposure of the perfused intestine to dissolved copper.

3.4.5 Effects of altering the gas composition on Ti uptake

Interestingly, the largest increase of net Ti uptake to the serosal compartment was caused by changing the gas mixture from 95% O_2 : 5% CO_2 to 99.5% O_2 : 0.5% CO_2 . It seems unlikely that the small 4% increase in O_2 content is physiologically important, since the tissue is already saturated with oxygen. The maximum amount of the oxygen in fresh water at air saturation (21% oxygen) is about 10 mg l⁻¹ (Eddy and Handy, 2012) and the gut tissue in the isolated perfused experiment was in direct contact with 99.5% of O_2 which means it is given at least five times the amount of the O_2 in fresh water. However, a change of CO_2 from 5 to 0.5% represents a profound change in acid-base balance for a freshwater fish (review, Goss et al., 1992). It is common practise in fish cell culture, and with perfused organ methods to use 95% O_2 : 5% CO_2 as a standard gas mixture for *in vitro* preparations. In this experiment, changing the gas mixture to 99.5% O_2 : 0.5% CO_2 (10 fold reductions in CO_2 content) caused a 13-fold increase in the initial Ti uptake rate in the TiO₂ NP exposed intestines (Table 3.7). A statistically significant effect was also noted for the bulk material, with a 3-fold increase in the initial Ti uptake with the 0.05% CO_2 gas mixture (Table 3.7).

This unexpected finding represents an important nano effect on bioavailability (10 fold elevation in Ti uptake with the nano treatment compared to the bulk treatment) that may also be of profound importance in real ecosystems because it is related to the partial pressure of gases in the external medium. The mechanisms involved require further investigation, but there are several possible explanations for this effect. One biological explanation is that the 5% CO₂ in the standard gas mixture for *in vitro* studies represents a severe acidosis for a freshwater fish (which normally have blood pCO_2 ten times lower than mammals, Eddy, 1977; Goss et al., 1992). Thus, lowering the CO₂ to

0.5% is simply moving it closer to *in vivo* values, and the Ti transport mechanism(s) are just recovering their normal (higher) transport rate following acidosis. This argument would work well for fish gill, but not the gut mucosa which can see some high CO₂ levels during digestive processes, and has sufficient carbonic anhydrase activity to convert CO₂ to bicarbonate (Grosell et al., 2009). In the present experimental conditions, the elevation of Ti uptake rate with decreasing CO₂ content of the gas mixture occurred with the expected changes of pH in the external medium associated with normal net base excretion in the intestine (> by about 0.4 pH units in the standard CO₂ mixture, and about 0.8 units in the 0.5% CO₂ mixture). The *p*CO₂ effect on Ti uptake can be explained by acid-base toxicity in the intestine.

An alternative explanation is that one of the mechanisms of Ti movement across serosal membrane is HCO_3^- -sensitive. Anion-dependent metal efflux pathways are known including a DIDS (4,4'-diisothiocyano stilbene-2,2'-disulfonic acid) and Cl⁻ sensitive Cu efflux in fish intestine (Handy et al., 2000), a Mg-anion transporter in mammalian cardiac muscle (Ödblom and Handy, 1999), and uptake of anionic Zn species on the Cl⁻/HCO₃⁻ exchanger in red blood cells (Torrubia and Garay, 1989). In the saline conditions of natural seawater Ti is almost exclusively present as TiO(OH)₂ (Van Den Berg et al., 1994), and water soluble forms of Ti readily form anion complexes with anions like citrate (Deng et al., 2007). It therefore seems theoretically possible that anion complexes of Ti could be exported to the blood on a basolaterally-located anion-exchanger, using the downward serosal to apical HCO₃⁻ gradient (low external CO₂) as a driving force.

The gas mixture effect is not easily explained by the chemical reactivity of TiO_2 with CO₂. Carbon dioxide gas spontaneously reacts on the surface of TiO_2 particles in the presence of alkali metals (e.g., Na⁺ K⁺, present in the gut at mmolar levels) to form carbonate ions (Krischok et al., 2002), that would be rapidly converted to HCO_3^- ions in

the gut lumen by carbonic anhydrase activity. The reaction on the crystal surface of the metal oxide normally uses oxygen from water molecules, but can utilise surface oxygen atoms from the TiO₂, especially where there are point defects in the crystal structure (Linsebigler et al., 1995). Thus a reaction of the crystal surface with CO_2 could liberate Ti ions as the oxygen atoms in the TiO_2 structure are used up. However, decreasing CO_2 in the medium would be expected to slow this reaction and therefore the availability of Ti ions in solution (less Ti dissolution). This is not consistent with the observed increase in Ti uptake at the lower CO₂ content (Fig. 3.8; Table 3.7). However, lowering the CO₂ would decrease external HCO_3^- generated by this reaction in the gut lumen and would drive some of the metal-anion transporters above. The gas mixture-dependent increase in Ti uptake was also observed with the bulk powder treatment, but to a lesser extent than the nano treatment. This is not necessarily a nano size effect, as numerous aspects of the crystal structure and spatial arrangement of particle surfaces will alter the CO₂ reaction outline above (Linsebigler et al., 1995), but clearly the gas mixture effect was stronger with nano compared to bulk material treatment in the experimental conditions used here.

3.4.6 Is there a material-type effect on tissue electrolytes in the perfused intestine?

There are some differences of Ti accumulation and uptake rates between bulk and nano treatments of TiO_2 in the perfused intestine (Fig. 3.7A and B; Table 3.7), but whether or not this translates into a nano effect of functional significance to the gut as an important osmoregulatory organ is less clear. Exposure to either material increased net water efflux across the intestine (Table 3.7, standard gas mix), but the bulk TiO_2 treatment caused a greater net water efflux than the nano treatment (Table 3.7) without dehydrating the tissue or altering tissue Na⁺ (Table 3.8). This suggests the bulk material was better at promoting physiological flux of water to the gut lumen, which would be an osmoregulatory advantage for a freshwater fish living in a very dilute environment

(Eddy, 2009). The bulk material also caused Mg^{2+} retention in the gut that was not observed in the nano treatment (Table 3.8), suggesting some material-specific effects on magnesium homeostasis. In the hind intestine, this was also accompanied by elevation of tissue Ca^{2+} due to treatment with the bulk material only (Table 3.8), implying some regional effect on calcium homeostasis. Similar observations were made for K⁺ in the mid intestine (Table 3.8). The depolarising effect of increased tissue K⁺ (i.e., more positive mucosal membrane potential) might also contribute to the difference in Ti uptake rate between the bulk and nano forms in this experiment (positive potential slowing Ti influx). The experiment with the 0.5% CO_2 gas mix also revealed a bulk material-specific elevation of tissue electrolytes compared to all other treatments, especially on tissue Na⁺ (Table 3.8). The reasons for this are unclear, but for example, bulk material interference with sodium carbonate co-transport is worthy of investigation.

Some mitochondrial damaged was found in the gut tissue after been exposed to both bulk and TiO₂ NP (Fig. 3.9 and 3.10). This damage included some degradation of the mitochondrial structure and swelling with the loss of definition of the cristae in the mitochondria. This observation is consistent with some oxidative injury of some of the mitochondria. The same mitochondrial damaged in the gut was shown in earthworms treated with 5 g kg⁻¹ of TiO₂ NPs (Hu et al., 2010), the authors reported disorganization, fracture and reduction (in size) of the mitochondria, and loss of the structure of the cristae. The physiological significance of these phenomena relates to the absorptive function of the epithelium. For example, during active trace element absorption from the food, trout intestinal epithelial cells show intact mitochondria which are aligned directly under the microvilli on the apical surface (Kamunde et al., 2003), apparently to provide energy for forming absorptive vesicles containing the micronutrients. Disrupted mitochondria would be less able to perform such functions.

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Conclusions

This study shows that the isolated perfused intestine preparation works with NPs, and the viability criteria remain good with these materials. Additionally, this work demonstrates that Ti from TiO₂ can cross the vertebrate intestine to appear in the serosal compartment, and there is a material-type effect with much faster net uptake of Ti from the nano form. Ti absorption from both bulk TiO₂ and nano form is time-dependent, saturable and the uptake mechanism is affected by the gas mixture used. All these factors suggest an active absorption mechanism happened. Therefore, from a regulatory perspective, a dietary exposure hazard to ecosystems and human health should be considered in risk assessments. The net uptake rates reported here for Ti are approximately 1-3 nmol g⁻¹ h⁻¹, identical to those for copper (Handy et al., 2000), but about 10 fold less than Hg uptake rates (40-50 nmol g^{-1} h⁻¹ for 0.2-2 mg l⁻¹ of Hg metal, Hoyle and Handy, 2005) in the perfused intestine. The oral uptake hazard for Ti is therefore of a similar magnitude to other metals (see also Ramsden et al., 2009). The fact that the nano form is taken up faster than bulk TiO₂ also implies that risk assessments based on the bulk material will underestimate the nano accumulation hazard. The assumption that the traditional bulk forms of TiO_2 powders are not appreciably absorbed via the gut should also be reconsidered. Notably, the gas mixture effect observed in the present study, adds a new dimension to hazard assessment not previously considered for metals or NMs. For ecosystems this could include considering changes in environmental pCO_2 associated with the exposure to TiO₂.

Chapter 4

Behaviour of TiO₂ NPs in the gut lumen and pharmacological studies on the uptake mechanism across the isolated perfused intestine of rainbow trout: nystatin and vanadate sensitive components

Abstract

There is limited information on the uptake mechanism of NPs and their pathway(s) across the gastrointestinal tract. In this study, different kinds of inhibitors (pharmacology treatments) were used to determine the pathway(s) for the uptake of Ti from TiO₂ exposures across the gut epithelium of the isolated gut perfusion of rainbow trout (Oncorhynchus mykiss). Pharmacological investigations of the uptake mechanism(s) of the Ti from TiO₂ NPs in perfusions gassed with 95% O₂: 5% CO_2 showed that addition of 10 mmol l⁻¹ potassium cyanide (KCN) to the mucosal and serosal solutions caused an elevated accumulation of Ti in the tissue, but this did not stop transepthelial uptake into the serosal compartment with 2.56 ± 0.65 nmol g⁻¹ h⁻¹ of Ti from the TiO₂ NPs (mean \pm S.E.M., n = 6). The addition of 100 µmol 1⁻¹ sodium orthovanadate, a P-type ATPase inhibitor, to the serosal solution increased the initial net uptake rate of Ti from TiO₂ NPs by 1.5 fold to 2.29 ± 0.64 nmol g⁻¹ h⁻¹ (mean \pm S.E.M., n = 6) compared to the TiO₂ NP treatment without vanadate. However, the overall net uptake in 4 h of Ti to the serosal compartment was reduced by 2.8 fold after vanadate additions. In contrast, adding 120 IU ml⁻¹ nystatin, an endocytosis inhibitor, to the mucosal solution completely blocked the uptake of Ti from both TiO₂ NPs and the bulk powder. The pharmacological responses with TiO_2 NPs showed some differences compared to their responses with bulk powder. We conclude that the uptake mechanism(s) involve a nystatin-sensitive endocytosis across the mucosal membrane.

4.1 Introduction

The precise mechanism(s) of NP uptake across the epithelia of fish (and mammals) remains controversial (Handy et al., 2008b; Shaw and Handy, 2011). Handy et al. (2008b) argues that apical entry of intact NPs into the epithelial cells of fish (gills, gut) is unlikely to occur via the traditional ion transport pathways (e.g., by diffusion through epithelial ion channels), because the NPs are simply far too large, but uptake into the cells by endocytosis remains possible. Similar arguments have been made for the mammalian gut (Panessa-Warren et al., 2006), and the cells of invertebrates (Moore, 2006). The diffusion of intact NPs through the paracellular route (e.g., via tight junctions) into the blood also seems unlikely (Handy et al., 2008b), due to the Ca^{2+} and Mg²⁺-rich microenvironment of the tight junctions which may lead the NPs forming aggregates. It seems more likely that most uptakes would be trans-cellular. However, it remains unknown whether any NPs taken up by putative endocytosis from the mucosal solution would be exported into the blood as intact particles or broken down to free metal ions. Information on the NPs transfer across epithelia is still unclear, relative to our knowledge on traditional dissolved metals (e.g., Foulkes and Bergman, 1993a; Handy et al., 2002a review).

The aim of this study was to pharmacologically investigate the uptake mechanisms of NPs using the isolated perfused intestine (Handy et al., 2000), and to also interpret any uptake in the context of particle behaviour in the gut lumen as well as surface bindings on the intestinal mucosa. The pharmacological investigations here involved using: (i) potassium cyanide (KCN) which is a general inhibitor of aerobic metabolism and often reduces the availability of ATP in cells for energy-dependent processes such as active ion uptake or movement of the cytoskeleton (endocytosis, cell volume control, etc.,); (ii) sodium orthovanadate which is a P-type ATPase inhibitor which non-specifically blocks the ATPase family of ion pumps including the Na⁺-K⁺

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ATPase, Ca^{2+} -ATPase, and Mg^{2+} -ATPase; (iii) nystatin, a putative inhibitor of the lipid raft-caveolae endocytosis pathway in animal cells. To interpret whether apparent uptake had any surface-bound components of Ti, some rapid solution dipping experiments were done to measure the instantaneous adsorption of TiO₂ to the intestine. In addition, and the possibility of free metal ion dissolution from the particles was explored using a dialysis method to aid interpretation of particle versus free metal ion absorption.

4.2 Methodology

4.2.1 Surface binding experiment

To aid data interpretation on Ti accumulation in the mucosa, some rapid solution dipping experiments were done to quantify surface binding of Ti. The approach followed a well-established method for fish tissue (Handy and Eddy, 2004) which is also well known for metal salts (Shaw and Handy, 2011). The technique involves allowing the tissue to instantaneous adsorb metal onto the surface of the epithelium over a few seconds (i.e., before true uptake can occur) and then determining the metal concentration on/in the tissue. Measurements were made on both mid and hind intestine. Briefly, fish were sacrificed and the intestinal tract was quickly removed, everted, and then rinsed in clean saline (see Chapter 3). Pieces of mid or hind intestine (n = 6 fish per treatment) were then dipped (mucosa facing the solution) in 500 ml of fresh physiological saline for 15 sec (control dip to equilibrate the surface of the tissue with the saline), then dipped in to 1 mg l^{-1} of TiO₂ NPs or bulk TiO₂ in 500 ml physiological saline for 30 sec. Finally, tissues were rinsed of excess TiO_2 by dipping in two consecutive 500 ml beakers of clean physiological saline (total of 15 seconds rinsing) (Fig. 4.1). Tissues were then digested in nitric acid and analysed for total Ti metal concentrations as described in Chapter 2.



Figure 4.1 Method of the surface binding tissue experiment showing the processing of the everted tissue dipping first in 500 ml of clean physiological saline for 15 sec then exposed the tissue to the 1 mg 1^{-1} of TiO₂ for up to 30 sec and finally dipped in same clean saline for 15 sec rinsing.

4.2.2 Dialysis experiments

These experiments were conducted to determine whether an apparent dissolved Ti metal fraction could be released from the TiO₂ NPs or the bulk material contained within dialysis tubing in the experimental conditions used for the gut perfusions. The rates of appearance of total apparently dissolved Ti (i.e., measured total Ti that could pass through the dialysis membrane into the external media) from the TiO_2 NPs or bulk TiO_2 in the physiological saline (described in Chapter 2) was determined using the standard gas mix and the low CO₂ gas mixture, at pH 7.4. In addition, a saline mimicking intracellular conditions (Handy et al., 1996) was also tested, which contained (in mmol 1⁻¹): KCl, 140; NaCl, 10; HEPES, 10 and adjusted to pH 7.2 to represent cytosolic intracellular pH. The intracellular saline was also made at pH 3, to determine whether an extremely acidic pH in an intracellular compartment (e.g., lysosomes) would cause the release of apparently dissolved Ti through the dialysis tubing. All glassware was acid washed (5% nitric acid) and triple rinsed in ultrapure Milli-Q water. The dialysis method is based on Handy et al. (1989) with modifications for NMs. Briefly, 8 ml of 100 mg l^{-1} of TiO₂ NPs, bulk TiO₂ stock dispersions made in the appropriate saline, or a control (the appropriate saline with no added TiO₂) were placed carefully into dialysis bags (7 cm long x 25 mm wide, dialysis tubing product code: D9777, cellulose membrane with a molecular weight cut off at 12,000 Da, Sigma-Aldrich, St. Louis, USA) and the ends closed with Medi-clips to prevent any leakage. The bags were immediately dialysed in a glass beaker containing 492 ml of the appropriate saline (bringing the total volume to 500 ml). Care was taken to use beakers of identical shape/size for the experiments, and each experiment was performed in triplicate (3) separate beakers) for each material and external medium (intracellular saline pH 7.2 and pH 3, physiological saline with standard gas mix and low gas mixture). The solutions in the beakers were gently agitated with a multipoint magnetic stirrer (RO 15P power, IkaWerke GmbH & Co. KG, Staufen, Germany) for 24 h at room temperature (21 °C). Samples of the external media (4.5 ml) were taken from each beaker at 0, 0.5, 1, 2, 3, 4, 6, 8 and 24 h, the pH of the samples was recorded at each time point. At the end of experiment, the remaining contents of the dialysis bag were also collected. All samples were then analysed by ICP-OES for total Ti metal concentration.

4.2.3 Preparation of the isolated perfused intestine

The preparation of isolated perfused intestine was performed exactly as described in Chapter 2 with some additions. Briefly, a series of perfusions were used to investigate the effects of inhibitors (n = 6 perfusions per treatment for each inhibitor). These included experiments with 10 mmol 1⁻¹ cyanide (KCN) added to both the perfusate and mucosal solutions to determine whether Ti uptake from exposure to TiO₂ NP had an energy-dependent component. Then in experiments using TiO₂ or bulk powder, the effects of adding 100 µmol 1⁻¹ sodium orthovanadate (Na₃Vo₄, an ATPase transporter inhibitor, Campbell et al., 1999; Handy et al., 2000) to the serosal solution was tested to explore whether ion transport was involved in Ti export from the gut cells to the blood. In separate experiments, additions of the endocytosis inhibitor, 120 IU ml⁻¹ nystatin (dose to produce 100% inhibition, Lewis et al., 1977), was added to the mucosal solution only.

The stock animals, preparations of the TiO_2 stock solutions and trace metal analysis, histology, calculations and statistical analysis were described in details in Chapter 2.

4.3 Results

4.3.1 Ti accumulation by surface binding experiment

Measurements from the rapid solution dipping protocol to determine surface-bound Ti on the mid and hind intestine revealed low concentrations of Ti metal. The total Ti metal concentration of dipped mid intestine was (means \pm S.E.M., n = 6); 0.007 \pm 0.001, 0.014 \pm 0.005 and 0.012 \pm 0.002 µmol g⁻¹ for control, bulk and TiO₂ NP exposures, respectively. The values for the TiO₂ treatments were not statistically different from the control (Kruskal-Wallis test, P = 0.47, Fig. 4.2), indicating negligible surface binding, and tissue concentrations were also significantly lower than those reported in the whole gut sacs or perfused intestine (*t*-tests, *P* values all < 0.05). Total Ti metal concentration in dipped hind intestine was (mean \pm S.E.M., n = 6); 0.006 \pm 0.001, 0.014 \pm 0.002 and 0.008 \pm 0.002 µmol g⁻¹ for control, bulk and TiO₂ NP exposures, respectively, with only the bulk being significantly higher than both control and NPs (ANOVA, *P* = 0.04, Fig. 4.2). Total Ti metal concentrations in the hind gut from the surface binding experiments were also significantly lower than the values reported in the whole gut sacs or intestinal perfusions (*t*-tests, *P* values all < 0.05), with the later representing (at the most) 27 and 26% of the total Ti reported for intact hind gut tissue in Table 3.1.



Figure 4.2 Total Ti metal concentrations in the tissues determined by ICP-OES in the dibbed mid (blue bars) and hind intestine (red bars) after exposed to 1 mg l⁻¹ TiO₂ (bulk or NP). Data are means \pm S.E.M. (n = 6 perfusions per treatment). * Statistically significant difference from control values (ANOVA or Kruskal-Wallis test, P < 0.05). # Statistically significant difference from bulk TiO₂ tretment (ANOVA or Kruskal-Wallis test, P < 0.05). There is no significant difference found between mid and hind intestine (*t*-test, P > 0.05).

4.3.2 Apparent total dissolved Ti from dialysis experiments with the mucosal solution and with intracellular salines

Dialysis experiments confirmed no measurable appearance of apparent total dissolved Ti metal from either the bulk or nano TiO₂ in the physiological saline used for the mucosal solution, indicating that the tissue was probably only exposed to particles. There were no measurable increases in the total Ti metal concentrations in any of the beakers from the dialysis experiments with physiological saline over 24 h (concentrations below detection limit, all < 1 μ g l⁻¹), regardless of the gas mixture (stand gas mix, or low CO₂ mix) or the type of particles added (the same finding for bulk and nano, data not shown). However, this was not the case for salines mimicking the intracellular environment.

Unlike the mucosal solution, the dialysis experiments using salines representing the intracellular ionic environment did show some appearance of measurable total Ti metal in the beakers from dialysis bags containing either the bulk or nano forms of TiO₂. In the pH 3 experiment (representing the low pH of vesicular compartments such as lysosomes) there was a small but detectable increase in apparent total dissolved Ti in the external media (beakers) after 3h in the TiO₂ treatments. Calculated cumulative Ti releases over 24 h were < 0.45 (values below detection in controls), 1.15, and 2.25 µg of Ti metal (mean of the triplicate measurement, absolute amount released) in to each dialysis bag for control, bulk and nano TiO₂, respectively. The rate of appearance of total metal in the external media in the beakers was much greater for the intracellular solution at pH 7.2 (i.e., representing the general cytosolic pH) with a clear time-dependent cumulative increase of the apparent total dissolved Ti concentrations in the beakers (Fig. 4.3). The maximum rates of appearance of cumulative total metal in the beakers (steepest part of the curves, not at equilibrium) were 9.7 and 6.9 µg Ti metal h⁻¹ (or 26 and 18 nmols Ti metal h⁻¹) for the bulk and nano TiO₂, respectively (control, at or

below detection limit). The beakers reached a steady-state maximum release of 46.0, and 34.7 μ g Ti metal over the 24 h for the bulk and TiO₂, respectively (Fig. 4.3).



Figure 4.3 The cumulative appearance of total Ti metal through dialysis tubing into the external medium of beakers for dialysis bags containing either bulk or nano TiO₂ using a saline mimicking the intracellular environment at pH 3 and pH 7.2. The data points are mean values derived from triplicate measurements (n = 3 beakers per treatment). Curves are fitted to the data points shown using a rectangular hyperbole function (one site, ligand binding to saturation) in SigmaPlot version 12. At pH 3, the bulk: y = 1.32*abs(x)/(2.51 + abs(x)), $r^2 = 0.96$. For TiO₂ NPs: y = 2.68*abs(x)/(3.90 + abs(x)), $r^2 = 0.97$. At pH 7.2, the bulk: y = 46.04*abs(x)/(2.815 + abs(x)), $r^2 = 0.99$, and TiO₂ NPs: y = 34.74*abs(x)/(3.126 + abs(x)), $r^2 = 0.99$. The controls were at or below detection limits, and so a curve is not fitted, but data points are shown for convenience (i.e., cumulative sum of detection limit values). The standard errors were too small to be visible on the figures.

4.3.3 Effects of inhibitors on the TiO₂ NP treatment

In general, the additions of inhibitors decreased the cumulative uptake of Ti to the serosal compartment, but without compromising perfusate flow in the preparations (examples in Fig. 4.4). This was confirmed for the TiO₂ NP treatment by a statistically significant decrease in the overall Ti net flux in the presence of 10 mmol 1^{-1} cyanide, 100 µmol 1^{-1} vanadate or 120 IU ml⁻¹ nystatin compared to the no added Ti control, or the bulk material alone (Kruskal-Wallis test, P < 0.05). Notably, nystatin completely abolished the Ti flux associated with the TiO₂ NP treatment. The effects of all these drugs on the TiO₂ NP treatment was accompanied by a statistically significant reduction in net water flux compared to the equivalent no-added drug TiO₂ treatment, but the water fluxes were still higher than the resting values from control intestines (Table 4.1) indicating that that additions of inhibitors did not reverse the effects of TiO₂ on net water efflux across the gut.

Despite some marked reductions in the net Ti flux associated with TiO₂ NPs exposure in the presence of inhibitors (Table 4.1), the effects of inhibitors on tissue total Ti metal concentrations during the TiO₂ NP exposures were modest. Serosal and mucosal applications of 10 mmol 1⁻¹ cyanide (KCN) did not stop Ti accumulation in the tissue for the TiO₂ NP treatment, but instead, increased the total Ti metal concentration in the tissue. Values were 0.08 ± 0.01 and $0.03 \pm 0.01 \mu$ mol g⁻¹ dry weight for the mid and hind intestine, respectively with cyanide (mean ± S.E.M., *n* = 6 for each, Fig. 4.5), being 4.1 and 1.4 fold higher than their respective tissue concentration in the presence of TiO₂ NPs without cyanide (statistically significant cyanide effect, *t*-tests, *p* = 0.001 and 0.035, respectively). For the TiO₂ NP treatment with vanadate (Fig. 4.5), some Ti accumulation in the tissue was observed in mid and hind intestine (total Ti metal concentrations in the tissue, mean ± S.E.M., *n* = 6; 0.033 ± 0.005 and 0.013 ± 0.004 µmol g⁻¹ dry weight, respectively), representing a small 1.7 fold increase in Ti in the

mid intestine compared to the TiO₂ treatment without the vanadate for the mid intestine (not statistically significant, *t*-test, P = 0.06), and a 0.4 fold decrease in the hind intestine (not statistically significant, *t*-test, P = 0.45). In the case of nystatin additions, the situation was similar with measurable Ti accumulation in the tissue (Fig. 4.5, 0.017 \pm 0.004 and 0.007 \pm 0.002 µmol g⁻¹ dry weight in the mid and hind intestine, respectively), but being only 10% and 68% lower than the respective values without the drug with TiO₂ NPs (*t*-tests, P = 0.26 and 0.39, mid and hind intestine, respectively).

Metabolic and ion transport inhibitors are expected to cause some dissipation of electrolyte gradients across tissues, all three inhibitors (cyanide, vanadate, and nystatin) caused statistically significant elevation of tissue Na⁺ and Ca²⁺ concentrations, and depletion of K⁺ in mid intestine, compared to the appropriate drug-free control (Table 4.2). For example the TiO₂ NP treatment with cyanide (mean \pm S.E.M., n = 6, mid intestine) showed Na⁺, Ca²⁺ and K⁺ concentrations of 405.2 \pm 54.7, 66.5 \pm 11.1 and 73.1 \pm 10.9 µmol g⁻¹ dry weight for each electrolyte, respectively. The same situation found in hind intestine, just for the K⁺ concentrations in vanadate and nystatin groups which appear significantly higher than TiO₂ NP drug free control. There were no drug effects on Mg²⁺ concentrations or tissue moisture content for both mid and hind intestine (Table 4.2).

4.3.4 Effects of inhibitors on the bulk TiO₂ treatment

The inhibitory effects of drugs on net Ti fluxes for the bulk material treatment were similar to those for the TiO_2 NPs (Table 4.1), with the drugs also abolishing or reducing the Ti net flux. There was a notable material-type effect on the vanadate-sensitive Ti flux (Table 4.1), with the Ti flux associated with the bulk material treatment being completely abolished, compared to only partial inhibition in the nano treatment (i.e., different sensitivities of the Ti flux associated with the bulk and nano treatments to vanadate). The total Ti metal concentrations in tissues following drug treatments in the

bulk TiO₂ exposure, showed statistically significant reductions in tissue Ti compared to the no-added drug Ti control (Fig. 4.5). For example, the addition of 100 µmol Γ^{1} vanadate to the perfusate solution showed total Ti metal concentrations associated with the bulk Ti treatment in the mid and hind intestine of 0.028 ± 0.006 and 0.007 ± 0.001 µmol g⁻¹dry weight of tissue, respectively (mean ± S.E.M., *n* = 6 for each), being 6.6 and 1.5 fold less than in the TiO₂ bulk treatment without drugs for mid and hind intestine, respectively (*t*-tests, *P* < 0.05). For the 120 IU ml⁻¹ nystatin, a similar effect on tissue total Ti metal concentrations was observed with Ti from the bulk Ti treatment with nystatin in the mid and hind intestine being 0.031 ± 0.004 and 0.008 ± 0.001 µmol g⁻¹dry weight of tissue, respectively (mean ± S.E.M., *n* = 6 for each), 6.3 and 1.4 fold less than that in the TiO₂ bulk treatment without drugs for mid and hind intestine, respectively (*t*-tests, *P* < 0.05). The tissue electrolyte response to inhibitors was similar to that described for the nano TiO₂ treatment above (Table 4.2), just for the Mg²⁺ concentrations with drug treatments which appear significantly lower than TiO₂ bulk drug free control for both mid and hind intestine (Table 4.2).

4.3.5 Histopathology

The histological examination of the perfusate intestine after 4 h perfusion with 1 mg 1^{-1} TiO₂ NPs or bulk using different inhibitors showed some minor injuries in the intestinal tissue (Fig. 4.6 and 4.7), the addition of 10 mmol 1^{-1} potassium cyanide to the serosal and mucosal solution showed some abnormal structure of the mucosal layer in compare to TiO₂ NP drug free control in five out of six animals. There were also some lesser effects with additions of 120 IU ml⁻¹ of nystatin to the mucosal solution in both bulk and nano TiO₂ which showed in three out of six animals per treatment. Serosal addition of 100 µmol 1^{-1} of sodium vanadate showed fusion of the occasional intestinal villi with TiO₂

NPs and shown in two out of six animals, but with the bulk TiO_2 the effects was minor,

just a slight appearance of lifting epithelium which showed in one out of six animals.



Figure 4.4 The cumulative appearance of total Ti metal in the serosal compartment (red bars, nmol Ti metal) of perfused intestine preparations from rainbow trout exposed to 1 mg Γ^1 of TiO₂ in the mucosal saline. Cumulative perfusate flow (blue diamonds, ml) is also shown. Graphs are representative examples of individual perfusions from replicated experiments (n = 6 perfusions per treatment; n = 7 perfusions for the control TiO₂ NP treatment), gassed with 95% O₂: 5% CO₂. (A) TiO₂ NPs as a control without drugs, (B) 10 mmol Γ^1 cyanide exposure with TiO₂ NP, (C) 100 µmol Γ^1 sodium vanadate exposure with TiO₂ NP, (D) 120 IU ml⁻¹ nystatin exposure with TiO₂ NP. Fish weights in the examples are 249.9, 360.7, 137 and 273.9 g, respectively.



Figure 4.5 Total Ti metal concentrations in the tissues determined by ICP-OES in the mid (blue bars) and hind intestine (red bars) of rainbow trout after 4 h perfusions with a mucosal side exposure to 1 mg 1^{-1} of bulk TiO₂ or TiO₂ nanoparticles (NP). The appropriate TiO₂ was added to the mucosal solution in the presence of the standard gas mixture (95% O₂: 5% CO₂), and in the presence of inhibitors: 120 IU ml⁻¹ nystatin (putative endocytosis inhibitor), 100 µmol 1^{-1} sodium vanadate (an ATPase inhibitor) and 10 mmol 1^{-1} cyanide (general inhibitor). Data are means ± S.E.M (n = 6 perfusions per treatment; n = 7 perfusions for the control TiO₂ NP treatment). * Statistically significant difference from the TiO₂ NPs drug free control (ANOVA or Kruskal-Wallis test, P < 0.05), # statistically significant difference from the bulk TiO₂ drug free control (ANOVA or Kruskal-Wallis test, P < 0.05). Letter (a) indicates statistically significant difference between mid and hind intestine (*t*-tests, P < 0.05).

Mucosal [TiO ₂] (1 mg l ⁻¹) and Drugs Concentration	Net Ti flux, $J_{\text{net,Ti}}$ (nmol g ⁻¹ h ⁻¹)		Net water flux, $J_{\text{net,H2O}}$ (ml g ⁻¹ h ⁻¹)		
	Initial rate	Overall rate	Initial rate	Overall rate	
Bulk TiO ₂	0.98 ± 0.47	0.85 ± 0.32	-26.28 ± 2.52	-28.54 ± 3.19	
TiO ₂ NP	$1.55 \pm 0.33^{\#}$	$2.38\pm0.68^{\#}$	$-17.56 \pm 4.00^{\#}$	$-18.01 \pm 4.16^{\#}$	
Bulk TiO ₂ vanadate	$< 0.004^{\#}$	$< 0.004^{\#}$	$-13.57 \pm 1.83^{\#}$	$-38.41 \pm 0.07^{\text{#a}}$	
Bulk TiO ₂ nystatin	$< 0.004^{\#}$	$< 0.004^{\#}$	$-09.97 \pm 2.05^{\#}$	-31.66 ± 0.08^{a}	
TiO ₂ NP cyanide	$2.56 \pm 0.65*$	$0.85\pm0.17*$	$-13.40 \pm 3.66*$	$-44.51 \pm 2.76^{*a}$	
TiO ₂ NP vanadate	$2.29 \pm 0.64*$	$0.83 \pm 0.22*$	$-16.26 \pm 2.66*$	$-49.17 \pm 3.88^{*a}$	
TiO ₂ NP nystatin	< 0.004*	< 0.004*	$-13.96 \pm 2.29*$	$-36.42 \pm 2.03^{*a}$	

Table 4.1 Effects of exposure to 1 mg l^{-1} TiO₂ with/without drugs on the Ti and water fluxes across the isolated perfused trout intestine.

Values are means \pm S.E.M. (n = 6 perfusions per treatment; n = 7 perfusions for the control TiO₂ NP treatment) expressed per gram dry mass of intestine per hour. Negative values indicate a net loss from the serosal solution, initial and overall rates data were calculated from cumulative perfusate data at 10 min and 4 h, respectively.* Statistically significant difference from TiO₂ NP drug free control values within columns (ANOVA or Kruskal-Wallis test, P < 0.05), # Statistically significant difference from bulk TiO₂ drug free control values within columns (ANOVA or Kruskal-Wallis test, P < 0.05). Letter (a) within rows indicates a statistically significant difference between initial and overall rates (*t*-test, P < 0.05). Drug concentrations were 120 IU ml⁻¹ nystatin, 10 mmol l⁻¹ cyanide, or 100 µmol l⁻¹ vanadate (see text for details).

Mucosal [TiO ₂]	Tissue [Metal] µmol g ⁻¹ dry mass						
$(1 \text{ mg } l^{-1})$	Na^+	K ⁺	Ca ²⁺	Mg^{2+}	Moisture %		
Mid Intestine							
Bulk TiO ₂	156.09 ± 24.33	212.02 ± 15.78	18.64 ± 3.45	20.15 ± 1.66	78.36 ± 2.14		
TiO ₂ NP	162.80 ± 34.59	$136.07 \pm 18.25^{\#}$	12.51 ± 1.34	$12.06 \pm 1.24^{\#}$	82.26 ± 1.72		
Bulk TiO ₂ vanadate	$258.92 \pm 13.52^{\#}$	$158.47 \pm 11.76^{\#}$	33.35 ± 6.50	$13.69 \pm 0.85^{\#}$	81.82 ± 1.23		
Bulk TiO ₂ nystatin	$253.28 \pm 15.29^{\#}$	$138.97 \pm 11.53^{\#}$	$36.69 \pm 8.16^{\#}$	$12.11 \pm 0.49^{\#}$	80.65 ± 1.38		
TiO ₂ NP cyanide	$405.20 \pm 54.74 *$	$73.09 \pm 10.96 *$	$66.48 \pm 11.06*$	10.87 ± 1.70	88.33 ± 1.09		
TiO ₂ NP vanadate	$427.10 \pm 50.53*$	160.08 ± 32.49	$56.90 \pm 10.23*$	17.17 ± 2.51	84.94 ± 2.61		
TiO ₂ NP nystatin	$229.45 \pm 21.40*$	$117.95 \pm 10.51 *$	$25.90\pm5.47*$	10.91 ± 0.86	84.54 ± 0.93		
Hind Intestine							
Bulk TiO ₂	116.48 ± 7.57	168.29 ± 5.93	18.00 ± 4.17	15.83 ± 0.69	81.54 ± 1.80		
TiO ₂ NP	98.65 ± 9.46	$121.03 \pm 7.05^{\#}$	$7.98 \pm 0.49^{\#}$	$9.99\pm0.59^{\#}$	80.85 ± 1.39		
Bulk TiO ₂ vanadate	$237.76 \pm 8.87^{\#}$	$164.60 \pm 4.42^{\#}$	13.23 ± 0.33	$13.49 \pm 0.28^{\#}$	82.08 ± 0.42		
Bulk TiO ₂ nystatin	$222.63 \pm 13.25^{\#}$	146.59 ± 5.74	15.30 ± 0.98	$11.50 \pm 0.46^{\#}$	80.65 ± 1.49		
TiO ₂ NP cyanide	$342.82 \pm 26.38*$	84.52 ± 8.99	$47.57 \pm 9.32*$	10.05 ± 0.53	88.11 ± 1.05		
TiO ₂ NP vanadate	$251.57 \pm 27.68*$	$174.39 \pm 18.22*$	$19.18 \pm 3.59*$	14.53 ± 1.29	82.02 ± 2.16		
TiO ₂ NP nystatin	$219.43 \pm 11.66*$	$156.27 \pm 7.17*$	13.93 ± 0.45	12.60 ± 0.48	85.24 ± 1.34		

Table 4.2 Total Na⁺, K⁺, Ca²⁺, and Mg²⁺ concentrations of gut tissue following exposure of isolated perfusate trout intestine to 1 mg l⁻¹ TiO₂ and different drugs for 4 h.

Values are means \pm S.E.M. (n = 6 perfusions per treatment; n = 7 perfusions for the control TiO₂ NP treatment) expressed as µmol g⁻¹ dry mass of intestinal tissue, except for moisture content (%) = ((wet weight – dry weight)/wet weight) x 100. * Statistically significant difference from TiO₂ NPs drug free control values within columns (ANOVA or Kruskal-Wallis test, P < 0.05), # Statistically significant difference from bulk TiO₂ drugs free control values within columns (ANOVA or Kruskal-Wallis test, P < 0.05).



Figure 4.6 Histology of the perfusate intestine after 4 h perfusion gassed with standard gas mix. (a) 1 mg Γ^1 TiO₂ NPs drug free control showing lifting of epithelial cell (L), (b) 1 mg Γ^1 TiO₂ NPs with 10 mmol Γ^1 potassium cyanide showing abnormal structure of the mucosal layer (in five out of six animals), (c) 1 mg Γ^1 TiO₂ NPs with 100 µmol Γ^1 sodium vanadate showing fusion of the intestinal villi (F), and appear of vacuoles (V) (in two out of six animals), (d) 1 mg Γ^1 TiO₂ NPs with 120 IU ml⁻¹ nystatin showing abnormal structure of the mucosal layer (in three out of six animals). Scale bar = 100 µm, sections were 8 µm thickness and stained with haematoxylin and eosin (n = 6 per treatment).



Figure 4.7 Histology of the perfusate intestine after 4 h perfusion gassed with standard gas mix. (a) 1 mg l⁻¹ bulk TiO₂ drug free control showing lifting of epithelial cell (L), (b) 1 mg l⁻¹ bulk TiO₂ with 100 μ mol l⁻¹ sodium vanadate showing lifting of epithelial cell (L) (in one out of six animals), (c) 1 mg l⁻¹ bulk TiO₂ with 120 IU ml⁻¹ nystatin showing abnormal structure of the mucosal layer (in three out of six animals). Scale bar = 100 μ m, sections were 8 μ m thickness and stained with haematoxylin and eosin (*n* = 6 per treatment).

4.4 Discussion

This is the first report on the uptake kinetics of NPs across the gut of rainbow trout using the isolated gut perfusion technique. Mechanism(s) uptake pathway of Ti from both bulk and nano forms of TiO_2 showed the effects of nystatin on the uptake, suggest at least part of the mucosal influx step is by endocytosis in the intestine. However, the effect of solute transport inhibitors also suggests a dissolved Ti component involved in Ti export from the tissue to the blood for bulk and nano TiO_2 . Further investigations using the rapid dipping solution experiment observed an apical uptake of the particles and the exposure is took place.

4.4.1 TiO₂ uptake through intestinal tissue

The both forms of TiO_2 showed that the Ti was accumulated in the mucosa, rather than the underlying muscularis (see Chapter 3, Table 3.2). The observed increases in total Ti metal concentrations in the mucosa are not easily explained by surface-binding (i.e., assumed adsorption) of TiO₂ (and/or Ti metal) to the exterior of the gut, with the rapid solution dipping experiments showing only about one fifth of the measured Ti being on the exterior within 30 sec (Fig. 4.2). There is usually some time dependence of surfacebinding in such experiments, but only for the first 90 sec (Handy and Eddy, 2004). A longer "dip" in the physiological saline of 60 sec might show slightly higher tissue Ti metal concentrations, suggesting more instantaneous binding of TiO₂ and/or Ti metal. However the instantaneous surface binding, of dissolved metals at least, is not linear over time with most of the binding occurring in the first 30 sec (Handy and Eddy, 2004) and a longer dip would at most add a further fifth (bring the total surface binding to 2/5th of the overall tissue metal concentration). This would still leave more than half the measured total Ti metal concentration representing internalised metal in the whole gut sac experiments. After 90 sec, radiotracer studies show that dissolved metals can be detected in the blood of trout (indicating transepithelial uptake, Handy and Eddy, 2004) and within 5 min the apparent accumulation is dominated by internal uptake of the metal. It would therefore seem logical that a 4 h exposure would give mostly internalisation (measured total Ti metal on the inside) rather than on the outside of the tissue.

4.4.2 Dissolution of Ti from both bulk and nano TiO_2 in different salines

The dialysis experiments showed that there was no appreciable dissolution of dissolved Ti metal from the mucosal saline in 24 h (gassed with 95% O₂: 5% CO₂ or 99.5% O₂: 0.5% CO₂), so the situation here is likely to be associated with the binding and settling of particulate TiO₂ with the secreted mucus. The notion of some small fraction of dissolved Ti ions in the cell is supported by the dialysis experiments, where the appearance of total Ti measured in the beaker outside the dialysis bag increased over time, but only when using the intracellular saline at pH 7.2 (Fig. 4.3B). This observation could be interpreted as a tiny fraction of dissolved Ti metal being released by dissolution from the particles, and subsequently diffusing through the dialysis membrane into the external media of the beaker. The rate of appearance of total metal in the external media in the beakers (18-26 nmol Ti metal h^{-1} depending on particle type) represents about 0.01% loss of metal from the particles every hour. If this interpretation is correct, this rate of release also represents the maximum dissolution rate of Ti metal from the particles in these specific experimental conditions. The dissolution of aqueous Ti metal species from TiO_2 particles has not been previously investigated in a high KCl media typical of the intracellular environment. However, Schmidt and Vogelsberger, (2009) recently investigated Ti dissolution from TiO₂ NPs (including P25 and anatase forms) in 0.1 mol l⁻¹ NaCl solution. These experiments were conducted over the whole pH range (pH 1-13) and for long durations (up to 1200 h), much longer than the 4 h perfusion here; but showed that TiO₂ will release metal ions by dissolution in NaCl solutions under certain conditions. At pH 3 or less, they detected dissolved Ti (around
0.1 and 1 μ mol 1⁻¹ Ti for pH 3 and pH 1, respectively). This observation tentatively suggests that TiO₂ NPs might also dissolve very slowly over several days in very acidic intracellular compartments (e.g., lysosomes). Schmidt and Vogelsberger, (2009) also measured the equilibrium solubility (at 500 h) of TiO₂ particles by absorptive stripping voltammetry, and showed that at pH 7, the P25 TiO₂ had a solubility of approximately 10⁻⁹ mol l⁻¹. This is a tiny amount of dissolved Ti from a physical chemistry perspective, but biologically, this would be a very relevant free ion concentration for metal homeostasis in cells. Schmidt and Vogelsberger, (2009) also calculated that this dissolved Ti would be present mainly as the dissolved Ti(OH)4 species at neutral pH, with the Gibbs free energy in favour of dissolution (overall solubility product, $10^{-9.05}$). There are some major differences between the dialysis experiments in this study and the work of Schmidt and Vogelsberger, (2009). The present study used much shorter durations (24 h for dialysis experiments) and so one might expect less dissolution of Ti in this experiments. However, KCl was also used instead of NaCl, and this seems to be important. The rate of release of apparent total dissolved Ti during dialysis was much faster in KCl solution (the intracellular saline, Fig. 4.3B) compared to the NaCl-rich mucosal saline. This may interpreted as an apparent difference in Ti dissolution rates between Na- and K-containing media. This needs further investigation, but may be due to the relative depolarising effect of K^+ or its greater diffusion coefficient (i.e., more mobile) compared to Na⁺ (Robinson and Stokes, 1968). Regardless, considering that trace metal homeostasis in cells operates at infinitely small free ion concentrations (e.g., Cu, maximally 10⁻¹⁸ moles cell⁻¹, Rae et al., 1999), the apparent total dissolved metal release rates in KCl-containing media here would be more than enough Ti release to enable Ti transport. Alternative explanations of the dialysis experiments, such as the NPs breaking up inside the dialysis bag (not observed) to release ~1 nm particles that might fit through the pore structure of the dialysis tubing cannot be completely excluded

in this experiment, but seems very unlikely given the KCl-effect above. It is also hard to imagine how primary particles of \sim 24 nm could weather to \sim 1-2 nm in 24 h without releasing significant quantities of metal ions. Notably, particles of a few nm wide were not observed in the electron micrographs of the original stock dispersions.

4.4.3 How does titanium associated with titanium dioxide particle exposures across the apical brush boarder membrane?

The discussion on events at the mucosal (apical) membrane of epithelia has focused on whether or not metallic NPs will be taken up directly across the apical membrane of epithelial cells as particles, and/or as dissolved metal (review, Shaw and Handy, 2011). In the present study the dialysis experiments showed that the appearance of total Ti metal in the mucosal solution was below the detection limits of ICP-OES, suggesting no measurable dissolution of dissolved Ti from the particles in the mucosal solution. The additions of nystatin (high dose, selected to completely inhibit endocytosis) caused about a 50 - 60% block of tissue Ti accumulation (depending on gut region and material, Fig. 4.5). Most of the apparent residue total Ti metal concentration in the tissue after nystatin treatment may simply represent the small amount of surface bound total Ti (regardless of form) detected in the solution dipping experiments (discussed above). Together, these observations argue in favour of some direct particle uptake at the mucosal membrane. However, the unstirred layer on the surface of epithelia is often a different chemical environment to that of the surrounding media (Handy and Eddy, 2004; Shaw and Handy, 2011) and the possibility of some dissolution of dissolved Ti metal from the particles in direct contact with the membrane cannot be excluded. Unfortunately the apical solute transport mechanisms for Ti metal are poorly understood, but the ionic radius of Ti^{2+} is about 1 Ångstrom (similar to Na^+ and Ca^{2+} ions, (crystaline values, 0.95 and 0.99, Robinson and Stokes, 1968). A theoretical possibility of facilitated diffusion through epithelial ion channels therefore exists (see Handy et al.,

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2002a). Facilitated diffusion would also be supported by the cyanide effect in the present study (Fig. 4.5), which did not prevent metal accumulation in the tissue, suggesting a passive uptake component across the mucosal membrane down the electrochemical gradient (as observed for Hg, Hoyle and Handy, 2005). A transferrinlike receptor mediated endocytosis (well known for Fe transport, see Bury and Handy, 2010), or non-specific uptake on divalent metal ion transporter 1 (Gunshin et al., 1997) are also theoretical possibilities. Furthermore, the reduction of net Ti uptake from TiO_2 NPs in the presence of cyanide (Fig. 4.4B) which was also sensitive to serosal additions of vanadate (Fig. 4.4C), suggests an energy-dependent efflux of Ti metal ions on a basolaterally-located ATPase into the blood as the most likely mechanism. There is no evidence in the literature of vanadate-sensitive ATPases being involved in exocytosis of particulate material from the gut mucosa to the blood at the serosal membrane. However, metal transporting ATPases are known to be involved in loading metal ions into vesicles. For example, it is well known that Cu ions are loaded into the Golgi compartment of cells via a Cu-ATPase (Dameron and Harrison, 1998), which is followed by subsequent vesicular trafficking of Cu to the serosal membrane (Handy et al., 2000). It remains possible that Ti could cross the cell using a similar ATPase-dependent trafficking system to the well-known Cu pathway, which is also known to be hijacked by other non-essential toxic metals like mercury (Hoyle and Handy, 2005). Tentatively, if particles are taken up into the tissue, it therefore implies that some dissolution of Ti from the materials was occurring in order to enable the vanadate-sensitive serosal efflux to occur. A separate nystatin-sensitive exocytosis pathway for particles to move from the epithelial cell to the blood is unlikely as nystatin completely abolished transepithelial uptake of both materials (e.g., Fig. 4.4D for TiO₂ NPs), but further research is needed on this aspect.

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4.4.4 Intestine tissue structure after drug exposures

Some injuries were observed in the gut tissue after exposed to different inhibitors, mainly shown in the mucosal layer, and without effects on the muscularis indicating the intestinal barrier was not compromised. The histological observations were limited, and with no clear TiO₂-dependent effects with the drugs present, and in only a few specimens. Potassium cyanide is considered as toxic via its interference with the electron transport train in mitochondria, preventing the tissue from using the oxygen (Eisler, 1991; Yamamoto, 1989). A localised subsequent hypoxia with metabolic acidosis could affect the structure of the mucosal layer. The distribution of the lipid rafts in the cell membranes could be altered by nystatin (Semis et al., 2012). It may, in theory, be possible for the membrane to be more vulnerable during lipid raft formation, and the oxidation potential of nystatin as a foreign substance has not been examined. Further work is needed to understand the effects of nystatin on the appearance of the mucosal layer in the intestine. For TiO₂ NPs, the vanadate treatment showed some vacuole formation with appearance of the occasional villi fusion. This was the same injury (but to a much lesser extent) to that observed by Federici et al. (2007), suggesting that the damage was caused by TiO_2 NPs exposure not by the vanadate. The bulk TiO_2 treatment showed a normal tissue appearance after exposure to vanadate with some minor lifting of epithelium (likely processing artefact) which appeared to be the same as the bulk TiO₂ (drug free control). Overall, this suggests no effects of vanadate on intestinal tissue integrity.

Conclusions

The present study indicates that the uptake of Ti from both bulk and nano TiO_2 had occurred across the intestine. This uptake was interpreted in the context of limited surface binding of Ti metal on the intestinal tissue. The dialysis investigation of the role of dissolved Ti ions tentatively suggests that some limited role in the uptake inside the cells. Further evidence of the mechanisms of uptake was found by testing inhibitors of the main uptake pathways in the gut; which indicated both nystatin-sensitive and vanadate-sensitive Ti uptake associated with TiO_2 exposures. This suggests that both particulate and dissolved metals components are involved in the overall uptake. This implies that environmental regulation for TiO_2 NPs cannot simply use the existing dissolved metal paradigm (Shaw and Handy, 2011), but a modification for particle uptake at the mucosal surface should be included in any biotic ligand model (BLM) for TiO_2 NPs. Further investigations on the mechanism of uptake are recommended using other inhibitors of endocytosis to determine, the sub-types of vesicular trafficking involved in the overall uptake.

Chapter 5

The uptake of different crystal structures of TiO₂ nanoparticles by Caco-2 intestinal cells

Abstract

The proposed uses of nanomaterials in food and different healthcare products heights the possibility of their uptake through the gastrointestinal tract in humans. The current study was applied to investigate the possible uptake of different forms of TiO₂ and their pathways for uptake into Caco-2 cells. Four different forms of $1 \text{ mg } 1^{-1} \text{ TiO}_2$ were used: (i) TiO₂ NPs as P25 with a composition of 75% anatase and 25% rutile, (ii) bulk TiO₂ with the same crystal structure as the P25, (iii) nano anatase TiO_2 and (iv) nano rutile TiO₂. The bulk TiO₂ showed the greatest uptake rate, being 14.1 nmol mg⁻¹ protein, while the nano rutile TiO₂ presents the lowest uptake rate, being 6.9 nmol mg⁻¹ protein. Another series of experiments investigated the mechanism of uptake of TiO₂ using two kinds of inhibitors, (i) 120 IU ml⁻¹ nystatin, an endocytosis pathway inhibitor and (ii) 100 µmol l⁻¹ sodium orthovanadate, a P-type ATPase inhibitor. Both vanadate and nystatin produced an increase in Ti accumulation from different forms of TiO₂. It is hypothesized that these drugs interfere with one of the endocytosis pathways. The results shows that 1 mg l^{-1} TiO₂ is actively taken up by Caco-2 cells with rates being influenced by the crystal structure and behaviour of TiO₂. The electrolyte concentrations in Caco-2 cells showed normal concentrations of Na⁺ and K⁺, but with an elevation in Mg^{2+} and Ca^{2+} concentration after exposure to different forms of TiO₂; suggesting specific material-type effects on electrolytes homeostasis

5.1 Introduction

Growing advances in nanotechnology have stimulated the development of many products in areas such as biomedicine, food, and cosmetics which may deliver chemicals or nutrients in a targeted and controlled manner to human beings. Titanium dioxide (TiO₂) is amongst the most widely used of metal oxide nanoparticles (NP) and is found in a variety of applications ranging from toothpastes to water treatment (Aitken et al., 2006; Fröhlich and Roblegg, 2012) and especially applications of TiO₂ NPs in cosmetics as transparent sunscreen blockers (Calzolai et al., 2012). Bulk TiO₂ is broadly used in food as a food colouring agent (normal white colour), food additives (E171) and in food packaging with daily ingestion estimated to be between 5 and 50 mg in the USA and Europe (Powell et al., 2010; Weir et al., 2012). According to Weir et al. (2012), 36% of the particles in E171 may be considered to be nano TiO₂ and its ingestion could have potentially toxic effects.

TiO₂ crystal structure and surface chemistry has been found to influence its toxicity (Chen and Mao, 2007). TiO₂ exists in three distinct forms brookite, anatase, and rutile, the latter two being more commonly produced (Chen and Mao, 2007). Studies have shown that anatase is more toxic than the rutile form with regards to its reactive oxygen species (ROS) promoting ability. This increased capacity for ROS generation compared to rutile TiO₂ has been shown to elicit cell damage and morphological changes (Sayes et al., 2006; Shi et al., 2007; Wang et al., 2008).

The pathways of inorganic NPs into and out of the cell membrane are still poorly understood. Dombu et al. (2010) suggested both endocytosis and exocytosis are involved in NP transport through epithelial cells. A study by Ryabachikova et al. (2010) examined the MDCK cells under electron microscope after exposure to different crystal forms of TiO_2 NPs. After 1 h, TEM images showed direct contact of NPs with the cell and NP filled and coated vesicles inside the membrane suggesting the possibility of receptor-mediated endocytosis (Ryabachikova et al., 2010). Only one study by Koenman et al. (2010) has been applied on Caco-2 human intestinal cell line exposure to TiO₂ NPs. Exposure of the cells to 10 or 100 μ g ml⁻¹ of TiO₂ NPs did not cause death to the cells or disruption of junctional complexes, TiO₂ appeared in the cells and underneath the cells suggesting that NPs transport through individual epithelial cells by transepithelial uptake is likely (Koenman et al., 2010). This has prompted research into which mechanisms are responsible for TiO₂ cell internalisation through the intestinal epithelium.

The overall aims of current study was to demonstrate the utility of the Caco-2 cell line for accumulation studies with TiO_2 NPs and then to compare the uptake of Ti from the bulk and nano forms (size effect), as well as the effects of different crystal structures (anatase and rutile). The approach included detailed pharmacological investigations of solute transport pathways for metals, as well as pharmacological studies to establish whether the different sizes or forms of TiO_2 were using different pathways to enter the cells. Finally, measurements of electrolytes in the cells, biochemical measurements on cell integrity, and electron microscopy investigations were made in the experiments to understand the physiological basis of any differences observed.

5.2 Methodology

Several experiments were performed using confluent monolayers of Caco-2 cells in culture medium (see below). The first series determined the Ti accumulation in confluent Caco-2 cells exposed to 1 mg l^{-1} of different forms of TiO₂ (bulk, nano, nano anatase or nano rutile) over 24 h, and also the effects of the TiO₂ exposure on cell viability (LDH release, cell morphology) and electrolyte concentrations. The second series of experiments involved pharmacological investigations to determine whether or

not the observed Ti accumulation involved either solute transport or endocytosisrelated pathways for uptake at the mucosal (apical) membrane. All data on Ti was normalized to nmol [Metal] mg⁻¹ cell protein (see below).

5.2.1 Cell culture

A human intestinal cell line, Caco-2 (brush border expressing, European collection of cell cultures) were routinely incubated in 75 cm², 200 ml flasks (Sterilin, Newport, UK) containing 15 ml of Dulbecco's modified eagle medium (DMEM) supplemented with 10% fetal bovine serum (FBS), 1% glutamine and 1% penicillin-streptomycin (100 IU Penicillin-100 µg ml⁻¹ Streptomycin), at 37 °C and gassed with 95% air: 5% CO₂, all the equipment (DMEM, FBS, glutamine and penicillin-streptomycin) was obtained from (Lonza, B-4800 Verviers, Belgium). For routine maintenance, the medium was changed every 48 h and the cells were sub-cultured by trypsination. Experiments were conducted on cells between passage 60 - 75 (cells were purchased at passage 45) and antibiotic were withdrawn to avoid possible interference with ion transport at least 2 passages before seeding the cells into 6-well plates (Sterilin, Newport, UK) for experiments. Preliminary trials were conducted to determine the optimum seeding density and time to confluence of the cells in 6-well plates by measuring the electrical resistance of the epithelium using an xCelligence real time label free cellular analysis system (Real-Time Cell Analyzer (RTCA), Roche Applied Science). In brief, the instrument monitors the electrical resistance of the cells grown on gold-coated wells, as cells adhere the resistance of the plate increases until confluence is reached, whereupon the relative resistance (cell index) remains constant. A seeding density of 5 x 10^4 cells/ cm² produced confluence by 24 h (Fig. 5.1). Subsequently for all experiments, cells were seeded at 5 x 10^4 cells/cm² in the 6-well plates and after an initial 24 h of growth, the cells were left for a further 48 h to ensure both 100% confluence and that the cells were rested. The cells were also visually inspected (Phase contrast microscope, Olympus/CK30-F200, Japan) each day until they became confluent, a description of general experimental methodology shown in Fig. 5.3 and 5.4. Cell viability was also checked by trypan blue exclusion, prior to seeding the flasks and 6-well plates, less than 90% viability was discarded. LDH was also checked in the external and internal cell cultures (see below) to measure the cell membrane integrity and the cytotoxic effects of different TiO₂ in compare to control LDH leak.



Figure 5.1 The cell index of Caco-2 cells after exposed to 1 mg l⁻¹ of different forms of TiO₂ showing the optimum seeding density of the cells at the 24 h of the exposure. Values are means \pm S.E.M (n = 4 plates per treatment).

5.2.2 Stock dispersions and materials characterisation

Four different types of TiO_2 were used (manufacturer's information): (i) bulk TiO_2 powder (ACROS, Titanium (IV) oxide, New Jersey, USA), with an unspecified mixture of anatase and rutile (but see below), with a purity of 98.0-100.5% TiO₂. (ii) Ultrafine TiO₂ NP type "Aeroxide" P25 (DeGussa AG, supplied by Lawrence Industries, Tamworth, UK) with a crystal structure of 25% rutile and 75% anatase TiO₂, purity was at least 99% TiO₂ (maximum impurity stated was 1% Si), the average particle size was 21 nm. (iii) Anatase TiO₂ NP form (US Research Nanomaterials, Inc, Houston, Texas, USA) with purity of 99% and 10-25 nm average particle size. (iv) Rutile TiO₂ NP form (US Research Nanomaterials, Inc, Houston, Texas, USA) with high purity, 99.9% and 30 nm average particle size. A stock solution of 500 mg l^{-1} for each type of TiO₂ was made (no solvents) by dispersing the particles in ultrapure Milli-Q water with manually vigorous shaking for 1-2 min. Preliminary studies identified the materials as source of bacterial infection to cell cultures (data not shown) and subsequently all stock dispersions and the dry powder were gamma irradiated to sterilise them (Red Perspex, Turntable Irradiation Geometry, Becton Dickinson, Belliver Industrial Estate, Plymouth, England). The radiation dose was 36.42 - > 40.72 kGy for 10 h to ensure sterility before starting cell culture experiments, and optimised to ensure the radiation did not interfere with the crystal structure of the materials.

The particle characterisation followed the similar protocol in Chapter 2. Briefly, the particle sizes were reported by using two methods. First, sub samples of the 500 mg I^{-1} stock dispersion were examined using transmission electron microscopy (TEM, JEOL-JEM.1400) (Fig. 5.2). The primary particle size measurement of 103.2 ± 16 nm (mean ± S.E.M., *n* = 7), 22.8 ± 0.6 nm (mean ± S.E.M., *n* = 169), 16.4 ± 2.4 nm (mean ± S.E.M., *n* = 6) and 30.8 ± 2.5 (mean ± S.E.M., *n* = 7) for bulk, P25, anatase and rutile, respectively, with the aggregates (viewed by electron microscopy) measuring 1162.8 ± 428.6 nm (mean \pm S.E.M., n = 6), 121.7 \pm 19.7 nm (mean \pm S.E.M., n = 8), 106.8 \pm 27.1 (mean \pm S.E.M., n = 6) and 115.1 \pm 17.9 (mean \pm S.E.M., n = 3), respectively.

The second method was to analyse the dispersion and distribution of TiO₂ in the medium by using nanoparticle tracking analysis (NTA, using a Nanosight LM 10, Nanosight, Salisbury, UK, laser output set at 30 mW at 640 nm). The media (DMEM) alone was filled with particles (29 nm smallest particle size and 156 nm mean values) so ultrapure Milli-Q water was used instead. TiO₂ was prepared by vigorous shaking instead of sonication and this may leave the stock full with aggregates particles, for this reason the concentration used here was 10 mg l⁻¹ to assess the measuring. The dispersion results gave a mean values of 179.3 ± 13.7, 7.1 ± 4.1, 142.3 ± 14.4 and 88.3 ± 34.1 nm (mean ± S.E.M., n = 3), for bulk, P25, anatase and rutile, respectively, While the mean of the smallest particles were 38.7 ± 3.5 , 7.5 ± 1.2 , 6.7 ± 1.2 and 7.7 ± 0.3 nm (mean ± S.E.M., n = 3), respectively (Fig. 5.2).



Figure 5.2 Electron microscope (EM) images showing (A) bulk TiO₂ particles (B) P25 (C) nano anatase and (D) nano rutile in 500 mg l⁻¹ stock solution (Milli-Q water). The images shows individual examples from triplicates (n = 7, 169, 6 and 7 for bulk, P25, anatase and rutile, respectively). Nanosight graphs showing the distribution of bulk TiO₂, P25, nano anatase and nano rutile in 10 mg l⁻¹ stock solution, graphs are individual examples from triplicate meaurements (n = 3).

5.2.3 Experiment 1: Time course of Ti accumulation from different forms of TiO_2

This experiment determined the time course of Ti accumulation from the different forms of TiO₂ in Caco-2 cells over 24 h. Confluent cells (72 h after seeding, as above) were exposed to the cell culture media (as above minus antibiotics) containing 1 mg Γ^1 of either no added TiO₂ (control), bulk, P25, anatase or rutile forms of TiO₂. Dosing of the wells was performed as follows: Media was made fresh on the day, TiO₂ (different forms) stock solutions were diluted with sterile distilled water from 500 mg Γ^1 to a concentration of 10 mg Γ^1 and then performing a 1: 10 dilution with cell culture media (1 ml stock: 9 ml media) to obtain a final concentration of 1 mg Γ^1 TiO₂ exposure. Cells were exposed for 24 h to each material, and then the cell mixture after scraped from the flasks was tested (see Fig. 5.3 and 5.4). Six plates per treatment at each time point: 0, 2, 4, 6, 8 and 24 h were analysed for Ti, electrolytes, LDH leak, and cell morphology (see below).



Figure 5.3 General experimental methods showing the processes of the experiment from seeding the cells till the analysis of samples. In all experiments, the cells were seeded in 6-well plates (two controls and different forms of TiO_2) and conducted in replicates (n = 6 plates per treatment). For the control, follow the same pattern of the experiment method with no exposure to TiO_2 .



SEM Examination

Figure 5.4 Experimental processing of the Caco-2 cells after been exposed the cells to 1 mg l^{-1} TiO₂ for 24 h. The cells were stained with Giemsa stain for the morphology examination (n = 3 plates per treatment). SEM grids of Caco-2 cells plate wells were obtained for the SEM examination (n = 3 plates per treatment). Caco-2 cells were fixed in a concentrated nitric acid and then analysed by the ICP-OES for the Ti and metal concentrations (n = 6 plates per treatment).

5.2.4 Experiment 2: The effect of nystatin and vanadate incubation on Ti accumulation

Having established the time course of Ti accumulation in Caco-2 cells from exposure to the different forms of TiO_2 the next stage was to assess the effect of mucosal (apical) additions of nystatin (a putative, endocytosis inhibitor, Iversen et al., 2011) and sodium orthovanadate (a P-type ATPase inhibitor, Cantley et al., 1978) on Ti accumulation. Cells were grown and dosed with 1 mg l^{-1} of the forms of TiO₂ as above, except that 1 h prior to dosing, cells were pre-incubated with either 120 IU ml⁻¹ of nystatin (dose to produce 100% inhibition, Lewis et al., 1977), or 100 µmol l⁻¹ sodium orthovanadate (enough to block metal transport in the intestine, Handy et al., 2000), compared to drugfree controls with and without added TiO₂. Both inhibitors were dissolved in 500 ml culture media (DMEM) to produce the appropriate concentration (see above) and preincubated for 1 h. This pre-incubation enabled the drugs to have direct contact with the cells without the risk of interference from the test materials (i.e., loss of bioavailable drug due to adsorption onto particles). The inhibitors remained in the media throughout the experiment, and the appropriate TiO_2 dose was simply added to the media after the initial 1h pre-incubation. Six plates per treatment for each inhibitor were conducted and after 24 h the cells was analysed as above.

5.2.5 Titanium determination and electrolytes in cells

In all experiments following TiO₂ exposure media was aspirated and centrifuged for 1 min (~160 g, Heraeus instruments, Biofuge *pico*, Germany) and collected for extracellular LDH analysis (see below). The cells were then collected to confirm Ti and/or electrolyte composition, the cells were washed twice with 2 ml of a sucrose washing buffer (300 mmol 1^{-1} sucrose, 0.1 mmol 1^{-1} ethylenediaminetetraacetic acid (EDTA), 20 mmol 1^{-1} 4-(2-hydroxyethyl)-1-piperazineethanesulfonic acid (HEPES), buffered to pH 7.4 with a few drops of 2 mol 1^{-1} Trizma base. Cells were scraped off the

dish (Fisher scientific cell scraper, 250 mm handle, 18 mm blade) and resuspended in 1 ml of a sucrose lysis buffer (the recipe above, but hypo-osmotic with only 30 mmol l^{-1} sucrose). The cell solution was sonicated (power 100 Watt, setting 8/ speed 22.5 kHz, Misonix incorporated, XL2000-010, New York) for 30 seconds to ensure the lysed sample was well mixed, centrifuged and a 200 µl aliquot fresh for LDH activity and protein determination (see below). For metal analysis, 1 ml of concentrated nitric acid was added to the remaining 800 μ l of the cell homogenate for total Ti, Na⁺, K⁺, Ca²⁺ and Mg²⁺ determination by (ICP-OES, Varian 725 ES). The standards used for ICP-OES calibrations were between 0-1, 0-20 and 0-200 mg l^{-1} for Ti, K⁺ and Na⁺, respectively and between 0-40 mg l^{-1} for both Ca²⁺ and Mg²⁺. All standards showed linear calibration with regression coefficient correlation of 0.99. To ensure acidity of the samples did not interfere with ICP- OES readout two sets of standards were made. One set were matrix matched and made in 30% nitric acid, the other made in 2% nitric acid. For the spike recovery test, samples (cell mixture) were spiked with 1 mg l^{-1} of different forms of TiO_2 and showed a good recovery of Ti metal measurement from the TiO_2 in different forms with 80.4 \pm 4.2, 82.2 \pm 1.5, 76.3 \pm 4.7 and 75.7 \pm 1.8% for bulk TiO₂, P25, anatse and rutile respectively (mean \pm S.E.M, n = 6 for each material type). Both sets of standards and spikes showed low coefficients of variation within and between samples (< 5%).

5.2.6 Lactate dehydrogenase and protein determination

Lactate dehydrogenase activity was measured in the cell culture media ("extracellular" LDH, n = 6 per treatment) and in the cell homogenates ("intracellular" LDH, n = 6 per treatment). For the former, 200 µl of cell culture media from each well was gently centrifuged for 1 min to remove any cell debris/turbidity (~160 g, Heraeus instruments, Biofuge *pico*, Germany) and 100 µl of the resulting supernatant was used in the LDH assay. For the cell homogenate, 200 µl were sonicated, centrifuged (see above) and then

100 µl of the resulting homogenate was used in the LDH assay using the method of Plummer, (1971), full description found in Chapter 2. LDH activity is expressed as µmol min⁻¹ ml⁻¹ of media, or µmol min⁻¹ mg⁻¹ cell protein as appropriate. For calculation of the % LDH leak from the cells, cells were lysed for total LDH (lysate + Media LDH) and then media LDH was calculated as a percentage of the total, % LDH leak = (100 X extracellular LDH) / (intracellular LDH).

LDH activity in cell homogenates was normalised for cell protein content, and protein was measured in triplicate using the bicinchoninic acid (BCA) method (MC155208, Pierce, Rockford, USA). The BCA reagent is a mixture of two reagents, reagent A, containing sodium carbonate, sodium bicarbonate, bicinchoninic acid and sodium tartrate in 0.1 M sodium hydroxide; and reagent B, containing 4% cupric sulphate. These two reagents were mixed in a ratio of 50:1 of reagent A: B. Briefly, 15 μ l of sample was added to 300 μ l of the BCA reagent and the absorbance read at 570 nm in 96-well plates (VERSA max, Molecular Devices, Berkshire, UK) against bovine serum albumin standards. Calibrations spiked with and without 1 mg l⁻¹ TiO₂ showed no interfere with the assay or colour reagent.

5.2.7 Cell morphology and scanning electron microscopy

Cell morphology was examined by light and electron microscopy during the cell culture, and prior to scraping the dishes at the end of the experiments. Briefly, cells were washed *in situ* on the cell culture dish with Dulbecco's phosphate buffered saline (DPBS) without calcium and magnesium (Lonza, B-4800 Verviers, Belgium), then fixed with 5 ml of 100% fresh methanol for 5 minutes, prior to staining with Giemsa (Giemsa's stain solution, IVD, England). Plates were examined wet using light microscopy (Nikon, 803923-Japan) and photographs taken using a digital camera (Pentax, K-X).

For scanning electron microscopy (SEM), the cells were fixed *in situ* on the culture plates and a disk cut out of the plate wells for mounting on SEM grids. Briefly, cells were washed in sucrose wash buffer (above) and fixed in 2.5% glutaldehyde in cacodylate buffer (0.2 M Sodium Cacodylate in water adjusted to a pH of 7.4) for 1 h. The cells were then washed twice in cacodylate buffer prior to being dehydrated in a graded series of ethanol. Cells were critically point dried and carbon coated (EMITECH-K850, K450X). Samples were imaged and elementally analysed (SEM, JEOL / JSM-7001F, Oxford Instruments INCA X-ray analysis system) using a 10 KV accelerating voltage, at a working distance of 10 mm.

5.2.8 Statistics

All data were statistically analysed used Stat Graphics Plus Version 5.1 and shown as means \pm S.E.M (standard error of mean), while Figures were drawn using Excel and Sigma Plot Version 12.0, full description in details found in Chapter 2. In the time course experiment, two-way ANOVA was used to determine treatment x time effects and the curve fitting for time-effects were done using Sigma Plot Version 12.0, and curves were fitted to the raw data, although graphs show the mean values for convenience.

5.3 Results

5.3.1 Cell health and viability

The control Caco-2 cells (not exposed to TiO₂ or treated with drugs) showed normal morphology during the experiments with the cells remained confluent and well attached to the dishes (Fig. 5.5a). This was also supported by the LDH activity which remained at around 0.1 IU ml⁻¹ or less of cumulative LDH release over the 24 h duration of the experiments (Fig. 5.6). The control Caco-2 cells also showed normal electrolyte levels with negligible variation in cell electrolyte concentrations. For example, in the time course experiment (triplicate plates/treatment) the electrolyte levels in control cells were (in nmol mg⁻¹ protein); 6859.2 ± 237.6 , 867.0 ± 26.3 , 16.0 ± 4.1 and, 48.0 ± 0.9 for Na⁺, K⁺, Ca²⁺ and Mg²⁺, respectively (mean \pm S.E.M, n = 6), representing a within experiment variation of 1 - 20% depending on the analyte. The controls were also reproducible across experiments (and batches of cells). For example, the control cells in the second experiment showed cumulative LDH values in the media of 0.1 IU ml⁻¹ or less, and the electrolyte concentrations in these controls were (in nmol mg^{-1} protein); 4518.6 \pm 160.9, 718.9 \pm 14.2, 10.6 \pm 1.2 and 42.6 \pm 1.1 for Na^+, K^+, Ca^{2+} and Mg^{2+} (mean \pm S.E.M, n = 6). Overall across all experiments, the between experiment variation for electrolyte concentrations in cells from the unexposed/untreated controls was 10 - 30% depending on the analyte, (sodium and calcium varying the most) with cumulative LDH activity in the media remaining low (0.1 IU ml^{-1} or less).

The morphology of the Caco-2 cells after incubating with different forms of TiO_2 over 24 h exposure showed normal cells with good confluent and well attached to the wells in compare to control (Fig. 5.5b, c, d and e). Cumulative LDH activity release in the external media over 24 h remained low with 0.1 IU ml⁻¹ or less for all different forms of TiO_2 (Fig. 5.6), the bulk showed the lowest LDH release with 0.04 IU ml⁻¹ in compare to other forms of TiO_2 . The percentage LDH leak through the Caco-2 cells

after exposed to different forms of TiO_2 was not significantly difference in compare to controls (ANOVA or Kruskal-Wallis test, P > 0.05, Table 5.1) indicating that the cell membrane integrity was not affected by different forms of TiO₂.



Figure 5.5 Morphology of the Caco-2 cells after 24 h exposure. (a) control (b) 1 mg l⁻¹ bulk TiO₂ (c) 1 mg l⁻¹ P25 (d) 1 mg l⁻¹ nano anatase (e) 1 mg l⁻¹ nano rutile showing normal morphology and well confluent of the cells in control and all treatment groups (n = 3 plates per treatment). Scale bar = 100 µm, stained with Giemsa's stain.



Figure 5.6 The effect of 1 mg l⁻¹ different forms of TiO₂ exposure on the LDH release in external media (extracellular LDH) over 24 h in Caco-2 cells. Values are means \pm S.E.M (*n* = 6). There were no statistically significant differences (ANOVA or Kruskal-Wallis test, *P* > 0.05) between groups (control, different forms of TiO₂).

	Total LDH activity (µmol min ⁻¹ ml ⁻¹)					
11021 mg 1	Control	Bulk TiO ₂	P25	Anatase	Rutile	
Extracellular LDH	0.05 ± 0.01	0.04 ± 0.02	0.09 ± 0.02	0.06 ± 0.03	0.06 ± 0.02	
Intracellular LDH	0.63 ± 0.02	0.61 ± 0.06	0.58 ± 0.07	0.56 ± 0.02	0.60 ± 0.03	
% LDH Leak	7.32 ± 1.67	6.57 ± 2.52	13.18 ± 1.70	9.52 ± 3.96	9.16 ± 3.07	

Table 5.1 LDH leak in Caco-2 cells over 24 h after incubation with different forms of TiO₂.

The effects of 1 mg l⁻¹ different forms of TiO₂ exposure on the total LDH activity release after 24 h in Caco-2 cells. Values are means \pm S.E.M. (n = 3-6). % LDH leak = (100 X extracellular LDH) / (intracellular LDH). There were no statistically significant differences (ANOVA or Kruskal-Wallis test, P > 0.05) between groups (control, different forms of TiO₂).

5. 3.2 Experiment 1: Time course of Ti accumulation from different forms of TiO₂

The time course of Ti accumulation for the different forms of TiO_2 are shown in Fig. 5.7, the unexposed control cells remained at a background level of around 2 nmol Ti mg⁻¹ cell proteins throughout the experiment. However, all the TiO₂ treatments showed a non-linear hyperbolic rise in the Ti concentration in the cells, which achieved steadystate concentrations by 24 h. This saturable rise in net Ti accumulation by the cells occurred without similar elevations of cell Na⁺ or K⁺ concentrations in all treatments. The Na⁺ and K⁺ concentrations remained between 558 - 668 and 779 - 814 nmol mg⁻¹ protein, respectively; indicating that the Ti response was Ti-specific and not an artefact of general electrolyte changes in the cells (not observed). There were some differences in the time courses of Ti accumulation for the different forms of TiO₂ (Fig. 5.7), and these were not explained by osmotic disturbances in the cells (no statistical differences by TiO₂ forms in the cell electrolyte contents, Table 5.2) or by differences in membrane permeability across the treatments. For the latter, the cumulative LDH leak by 24 h remained low at around 0.1 IU ml⁻¹ (not statistically different from the control, and no material-type effect, ANOVA, P = 0.066 Table 5.1). The net Ti accumulation by 24 h was in the following order by material type: bulk, P25, anatase, rutile and unexposed controls. The bulk material and P25 showed the largest net Ti accumulation in 24 h, saturating at 14.4 \pm 1.7 and 12.9 \pm 0.4 nmol mg⁻¹ protein respectively, but were not statistically different from each other (*t*-test, P = 0.473). The anatase form showed lower values with approximately 40% less Ti accumulation than either the P25 or bulk material (statistically different by 24 h, Kruskal Wallis test, P = 0.004). There were also some differences in Ti accumulation from the pure anatase and rutile forms, with the latter showing saturation at a slightly lower Ti concentration (Fig. 5.7). However, there was no statically significantly different showed between the anatase and rutile at 24 h (ttest, P = 0.295). The initial Ti accumulation rates calculated from the curves followed a similar pattern with Ti accumulation from the bulk and rutile being faster than those from anatase or P25 (Fig. 5.7) and were 5.3, 3.73, 3.58 and 4.48 nmol mg⁻¹ protein hr⁻¹ for bulk, P25, anatase and rutile, respectively. The 50% saturation of Ti from different forms of TiO₂ were 7.1 nmol mg⁻¹ protein at 1 h and 48 min, 6.82 nmol mg⁻¹ protein at 2 h and 42 min, 4.26 nmol mg⁻¹ protein at 1 h and 30 min and 3.47 nmol mg⁻¹ protein at 30 min for bulk, P25, anatase and rutile, respectively. The plateau for the Ti accumulation from both bulk and P25 were at 8 h with 13.0 and 11.6 nmol mg⁻¹ protein, respectively. Anatase was plateauing at 4 h with 6.78 nmol mg⁻¹ protein while the rutile was the most rapid one which plateauing at 2 h and 30 min with 6.08 nmol mg⁻¹ protein. At 24 h, the Ti from all different forms of TiO₂ exhibited statistically significant accumulation relative to control values (ANOVA or Kruskal-Wallis, P < 0.05) demonstrated a saturable time dependent response to 1 mg l⁻¹ different forms of TiO₂.



Figure 5.7 Titanium uptake in Caco-2 cells incubated with 1 mg l⁻¹ different forms of TiO₂ for 24 h. The data points are mean values derived from triplicate measurements (n = 6 plates per treatment for each time point) and are expressed as nmol [Ti] mg⁻¹ protein. Curves are fitted to the data points shown using hyperbolic exponential rise to the maximum (single, 3 parameters) in SigmaPlot version 12. The bulk: $y = 2.65+11.54*(1-\exp(-0.26*x))$, $r^2 = 0.90$, P25: $y = 1.16+12.54*(1-\exp(-0.22*x))$, $r^2 = 0.86$, anatase: $y = 1.71+7.01*(1-\exp(-0.30*x))$, $r^2 = 0.95$ and rutile: $y = 2.07+4.86*(1-\exp(-0.67*x))$, $r^2 = 0.73$. * Statistically significant difference from control values (ANOVA, P < 0.05). The control values were significantly lower than the treatment forms and so the curve is not fitted, but data points are shown for convenience.

5.3.3 Effects of TiO₂ exposure on intracellular electrolytes over 24 h

Caco-2 cells after exposed to 1 mg l⁻¹ of different forms of TiO₂ over 24 h exhibited different effects on cell electrolytes. Intracellular potassium significantly decrease in all treatments groups relative to controls (ANOVA, P = 0.001, Table 5.2). The lowest value of K⁺ was seen after rutile exposure, which dropped by 107 nmol mg⁻¹ protein compared to the control.

Intracellular magnesium significantly increased in all treatments groups in comparison to the control (ANOVA, P = 0.0003, see Table 5.2). P25 caused the highest effects on Mg²⁺ levels relative to other treatment groups, showing 10.8 nmol mg⁻¹ protein increases in comparison to the control magnesium level.

Calcium levels rose significantly in cells exposed to different forms of TiO₂ in comparison to the control (Kruskal-Wallis test, P = 0.04, Table 5.2). P25 caused the largest increase in intracellular calcium with a 24 nmol mg⁻¹ protein rise compared to the control. The exposure of different forms of TiO₂ had no statistically significant effects on intracellular sodium levels relative to the control (Kruskal-Wallis test, P =0.30, Table 5.2). However, changes in electrolyte levels were against the electrochemical gradient indicating that the membrane remained operative and did not passively leak electrolytes throughout the duration of the experiments.

5.3.4 Experiment 2: The effect of nystatin and vanadate incubation on Ti accumulation

Caco-2 cells were pre-incubated with 120 IU ml⁻¹ nystatin to establish endocytosis inhibition or 100 μ mol l⁻¹ vanadate (a P-type ATPase inhibitor) to enhance blocking of any Ti metal transport through intestine. Nystatin and vanadate treatments showed significantly increased levels of bulk Ti accumulation compared to the control (ANOVA, P = 0.0002, Fig. 5.8). Bulk Ti accumulation increased by 3 nmol mg⁻¹ protein compared to the control (10.76 nmol mg⁻¹ protein) when incubated with nystatin, reaching a final concentration of 13.76 nmol mg⁻¹ protein at 24 h. Incubating with vanadate further increased bulk Ti accumulation to 20.10 nmol mg⁻¹ protein at 24 h, an increase of 10 nmol mg⁻¹ protein compared to the control.

Nystatin caused an increase in P25 Ti accumulation (14.36 nmol mg⁻¹ protein), 4 nmol mg⁻¹ greater than the control (9.89 nmol mg⁻¹ protein). Incubating with vanadate caused additional Ti accumulation reaching a final concentration of 22.26 nmol mg⁻¹ protein, an increase of 12 nmol mg⁻¹ protein compared to the control with a significant increase for both drugs (Kruskal-Wallis test, P = 0.0005, Fig. 5.8).

Anatase Ti accumulation were significantly increased when the cells were incubated with nystatin or vanadate (Kruskal-Wallis test, P = 0.0015, Fig. 5.8). Incubating with nystatin caused a 2 nmol mg⁻¹ protein increase in accumulation compared to the control (9.47 nmol mg⁻¹ protein). Incubating with vanadate further increased anatase Ti accumulation to 22.04 nmol mg⁻¹ protein, an increase of 12 nmol mg⁻¹ protein compared to the control.

Rutile Ti accumualation did not follow the same pattern in comparison with the other forms of TiO₂. Nystatin incubation showed a significant increase in Ti accumulation from exposure to the rutile form (ANOVA, P = 0.004, Fig. 5.8), by a 4 nmol mg⁻¹ protein increase in compare to the control (5.38 nmol mg⁻¹ protein).

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Vanadate caused just a 0.9 nmol mg⁻¹ increases in Ti accumulation, however this was not deemed statistically significant in comparison to the control (*t*-test, P = 0.1602).



Figure 5.8 Drug effects on Ti accumulation in Caco-2 cells incubated with 1 mg l⁻¹ different forms of TiO₂ over 24 h. Data are means \pm S.E.M (n = 6) and expressed as nmol [Ti] mg⁻¹ protein. * statistically significant difference from the no drug control values (ANOVA or Kruskal-Wallis test, P < 0.05). # statistically significant difference from the nystatin values (ANOVA or Kruskal-Wallis test, P < 0.05).

5.3.5 Effect of nystatin and vanadate exposure on intracellular electrolytes over 24 h

The electrolyte levels in Caco-2 cells varied in response to drug incubation and remained consistent within TiO₂ treatments. Electrolyte levels in nystatin controls were $6303.7 \pm 468.1, 927.9 \pm 50.9, 25.7 \pm 9.5$ and 56.2 ± 2.9 nmol mg⁻¹ protein for Na⁺, K⁺, Ca^{2+} and Mg^{2+} , respectively (mean \pm S.E.M, n = 6). Additions of different forms of TiO₂ to Caco-2 cells incubated with 120 IU ml⁻¹ nystatin had no effects on intracellular Na^+ , K^+ and Mg^{2+} levels relative to the no TiO₂ positive control (ANOVA or Kruskal-Wallis test, P > 0.05, Table 5.2). However, Ca²⁺ values were significantly decreased in comparison to the no TiO₂ positive control (Kruskal-Wallis, P = 0.04). Cells incubated with nystatin exhibited a significant increase in intracellular potassium with all the different forms of TiO₂ relative to the no drug controls (ANOVA or Kruskal-Wallis test, P < 0.05, Table 5.2) and similar to the equivlaent exposures without drugs. The level of K^+ decreased with all forms of TiO₂ in comparison to control values with nystatin present (see Table 5.2). Intracellular magnesium showed significantly increased levels for all the different forms of TiO₂ relative to their no drug controls (ANOVA or Kruskal-Wallis test, P < 0.05), except in cells exposed to P25 which were found not to be significantly different from the no drug control. Nystatin had no effects on intracellular calcium or sodium levels in compare to the no drug control (ANOVA or Kruskal-Wallis test, P > 0.05).

Electrolytes levels in the vanadate controls showed lower values than the nystatin controls (above) being 3632.0 ± 427.1 , 466.9 ± 43.8 , 20.9 ± 3.6 and 30.9 ± 3.4 nmol mg⁻¹ protein for Na⁺, K⁺, Ca²⁺ and Mg²⁺, respectively in vanadate-treated cells (mean \pm S.E.M, n = 6). K⁺ and Mg²⁺ levels in all TiO₂ forms incubated with vanadate showed a significantly lower values relative to the nystatin in all different forms of TiO₂ (ANOVA or Kruskal-Wallis test, P < 0.05). Caco-2 cells incubated with 100 µmol 1⁻¹ vanadate and different forms of TiO₂ displayed significantly decreased levels of

intracellular potassium in all TiO₂ forms relative to the no drug controls (ANOVA or Kruskal-Wallis test, P < 0.05), but there were no effects found on potassium levels in all TiO₂ forms relative to the no TiO₂ positive control (ANOVA, P = 0.91, Table 5.2). Cells incubated with vanadate cause a significant decrease in intracellular magnesium in all different forms of TiO₂ relative to the no drug controls (ANOVA or Kruskal-Wallis test, P < 0.05), but showed a significantly increase of Mg²⁺ level in the anatase and rutile groups comparing to the no TiO₂ positive control (ANOVA, P = 0.047). Calcium levels increased significantly in cells after incubated with vanadate in both bulk and rutile groups relative to the no drug controls (ANOVA or Kruskal-Wallis test, P < 0.05). There was no effects on calcium levels in all different forms of TiO₂ control (ANOVA, P = 0.48). For the intracellular sodium levels, there was no significant effect found in compare to no drug control or no TiO₂ control (ANOVA or Kruskal-Wallis test, P > 0.05).

$TiO_2 1 mg l^{-1}$		Electrolytes (nmol mg ⁻¹ protein)					
		Na ⁺	\mathbf{K}^+	Ca ²⁺	Mg ²⁺		
No drugs	Control	4518 ± 160	718.8 ± 14.2	10.6 ± 1.2	42.6 ± 1.1		
	Bulk TiO ₂	4666 ± 576	$663.9 \pm 28.3*$	14.3 ± 7.0	47.8 ± 2.2		
	P25	4793 ± 231	$643.8 \pm 19.6 *$	$28.0 \pm 12.4*$	$53.4\pm2.7*$		
	Anatase	5109 ± 624	$637.9 \pm 27.6*$	$20.8\pm5.4*$	$49.8 \pm 1.4 *$		
	Rutile	4319 ± 350	$611.1 \pm 11.4*$	$19.1 \pm 8.2*$	$50.7 \pm 1.5 *$		
Nystatin	Control	$6303 \pm 468^{\#}$	$927.9 \pm 50.8^{\#}$	$25.7 \pm 9.4^{\#}$	$56.2 \pm 2.9^{\#}$		
	Bulk TiO ₂	5253 ± 445	$828.8 \pm 62.3^{\#}$	$19.8 \pm 2.2^{*^{\#}}$	$55.3\pm2.8^{\#}$		
	P25	5144 ± 308	$815.7 \pm 30.0^{\#}$	27.8 ± 5.7	55.0 ± 1.8		
	Anatase	5748 ± 455	$822.7 \pm 57.9^{\#}$	$14.4\pm4.6^*$	$56.2\pm4.4^{\#}$		
	Rutile	5202 ± 417	$807.1 \pm 41.2^{\#}$	$17.7 \pm 3.1*$	$55.0 \pm 3.1^{\#}$		
Vanadate	Control	3632 ± 427 †	466.8±43.8†~~	$20.9\pm3.6^{\sim}$	30.9 ± 3.4 †~		
	Bulk TiO ₂	4246 ± 406	479.6 ± 36.9 †~	$25.2\pm4.6^{\sim}$	38.9 ± 3.1 †~		
	P25	3982 ± 311	469.3 ± 26.9	18.0 ± 7.2	39.4 ± 2.7 †~		
	Anatase	4457 ± 216 †	513.3 ± 16.9 †~	18.7 ± 4.1	43.8 ± 2.3 †*~		
	Rutile	4534 ± 568	490.7 ± 37.1 †~	$29.1 \pm 2.4^{\sim}$	40.9 ± 2.8 †*~		

Table 5.2 Total Na⁺, K⁺, Ca²⁺ and Mg²⁺ nmol [Metal] mg⁻¹ protein concentration in Caco-2 cells after exposed to 1mg l⁻¹ different forms of TiO₂ over 24 h.

Data are expressed as means \pm S.E.M (n = 6 for each group). * statistically significant difference from the control values within columns, treatments and electrolytes (ANOVA or Kruskal-Wallis test, P < 0.05). # statistically significant difference of nystatin values relative to the no drug trials within columns, treatments and electrolytes (ANOVA or Kruskal-Wallis test, P < 0.05). ~ statistically significant difference of vanadate values relative to the no drug trials within columns, treatments and electrolytes (ANOVA or Kruskal-Wallis test, P < 0.05). ~ statistically significant difference of vanadate values relative to the no drug trials within columns, treatments and electrolytes (ANOVA or Kruskal-Wallis test, P < 0.05). † statistically significant difference of vanadate values relative to nystatin values within columns, treatments and electrolytes (ANOVA or Kruskal-Wallis test, P < 0.05). † statistically significant difference of vanadate values relative to nystatin values within columns, treatments and electrolytes (ANOVA or Kruskal-Wallis test, P < 0.05).

5.3.6 Presence of particles within the Caco-2 cells

Scanning electron microscopy (SEM) of the Caco-2 cells was employed to investigate the presence, condition and organization of microvilli on the apical surface of control cells compared to cells exposed to the different forms of TiO₂ (Fig.5.9), and also to explore the presence of the TiO_2 particles in/on the cells by using the X-ray microanalysis detector (Fig. 5.10 and 5.11). Control cells showed good microvilli growing on the surface of the cultured cells after 96 h incubation. The exposure of Caco-2 cells to 1 mg l^{-1} of different forms of TiO₂ for 24 h showed different effects on the microvilli growth; bulk and anatase have a good development of the microvilli growth which showed similar organization to control cells. P25 and rutile seems to have the lowest microvilli growth density with the decreased numbers of the microvilli appearance on the apical surface of the cells relative to the no treated control cells (Fig. 5.9). For all TiO_2 forms, the Ti was detected by pinpoint X-ray analysis which observed the Ti particles in/on the Caco-2 cells and appear to be penetrate the cells and made a contact with the cell membrane (Fig. 5.10), some of the particles are suggest to be translocated in the nucleus (Fig. 5.11d-f) and other particles have been shown on the cell surface (Fig. 5.11a-c).


Figure 5.9 Scanning electron microscopy of Caco-2 cells after 96 h incubation showing the microvilli growth density on the apical surface of the cells after exposed to 1 mg l⁻¹ different forms of TiO₂ for 24 h. (a) control, (b) bulk, (c) P25, (d) anatase and (e) rutile. Scale bar = 1 μ m (*n* = 3 per treatment).

Chapter 5



Figure 5.10 Scanning electron microscopy of Caco-2 cells after 96 h incubation showing presence of the Ti particles underneath the cells after exposed to 1 mg l⁻¹ different forms of TiO₂ for 24 h (n = 3 per treatment). (a) Secondary electron image (SEI) of the bulk cell surface, scale bar = 1 µm; (b) Backscatter image (BSI) of 'a' image showing subsurface Ti with xrma spectra; (c) SEI of the P25 cell surface, scale bar = 1 µm; (d) BSI of 'c' image; (e) SEI of the anatase cell surface, scale bar = 1 µm; (f) BSI of 'e' image, (g) SEI of the rutile cell surface, scale bar = 100 nm; (h) BSI of 'g' image.



Figure 5.11 Scanning electron microscopy of Caco-2 cells after 96 h incubation showing presence of the Ti particles on the surface of the cells after exposed to 1 mg l⁻¹ different forms of TiO₂ for 24 h (n = 3 per treatment). (a) Surface bound of P25 with spectra, scale bar = 3 µm; (b) Surface bound anatase with associated spectra, scale bar = 2 µm; (c) Surface bound rutile with associated spectra, scale bar = 2 µm; (d) SEI image of the cell surface exposed to P25. The yoke like structure in the centre is the nucleus, scale bar = 10 µm; (e) SEI image of the cell surface exposed to rutile showing the nucleus in the centre, scale bar = 1 µm; (f) A high magnification BSI of 'e' highlighting a Ti dense region within the left side of the nucleus.

5.4 Discussion

This experiment investigated the processes involved in the uptake and accumulation of different forms of TiO_2 using Caco-2 cells. The main findings were that Ti uptake rates from bulk TiO_2 were greater than that from the different nano forms of TiO_2 , which indicated a material-type effect towards uptake of TiO_2 in Caco-2 cells. Following nystatin (an endocytosis inhibitor) and vanadate (a P-type ATPase inhibitor) incubation, an explanation of the data is that the mechanisms of TiO_2 uptake are likely mediated by endocytic mechanisms. Furthermore, exposure to different forms of TiO_2 (bulk and nano) altered cell electrolyte homeostasis.

5.4.1 Cell health and viability

The morphology of Caco-2 cells in the control and in all forms of TiO₂ was good in general, with normal morphology and well-confluent cell appearance (Fig. 5.5). There was no effect on the cell structure after exposure to different forms of TiO₂ in comparison with controls. The present findings seem to be consistent with other research which found no alteration in the morphology or density of the Caco-2 cells after exposure to 100 μ g ml⁻¹ TiO₂ NPs for 4 h, 24 h and 72 h compared to control cells (Fisichella et al., 2012).

The LDH activity was performed to ensure that the cell membrane integrity had been preserved throughout the experiments. There was no LDH leakage shown in either the controls or after exposing the cells to 1 mg 1^{-1} of different forms of TiO₂. Control Caco-2 cells showed normal electrolyte levels, with negligible variation in electrolyte concentrations after exposure to different forms of TiO₂. Taken together, the normal morphology of the cells, negligible LDH leak, and normal electrolyte concentrations in Caco-2 cells indicate that cells were in good viability and have the utility for measuring the accumulation of Ti from different forms of TiO₂.

5.4.2 Uptake and accumulation of Ti from different forms of TiO₂ by Caco-2 cells

Ti uptake from different forms of TiO_2 in Caco-2 cells showed a steady, saturable, time dependent appearance over the 24 h experiment. This most likely indicates that Ti uptake involves active transport by endocytosis mechanism(s), which are influenced by particle characteristics, resulting in the nonlinear (saturable) uptake curves observed (Fig. 5.7), so the passive diffusion through the cell membrane or leakage of dissolved Ti seems unlikely. This is also indicated by Brun et al. (2011), who argue that no partial dissolution occurs for NPs after the Ti is internalised in Caco-2 cells as the nanoparticulate state. However, details of their solution preparation, exposure conditions, replication and controls were not fully documented in this preliminary conference report. Fröhlich and Roblegg (2012) also suggested that endocytosis is the most likely mechanism for the uptake of NP into intestinal epithelial cells.

The net uptake rate of Ti from bulk TiO₂ in Caco-2 cells was greater than the other NP forms of TiO₂. A possible explanation for this might be that the larger particles of the TiO₂ remained within the submucosa of the intestine and colon (Jani et al., 1990) or this might be related to the behaviour of bulk TiO₂ in the culture media. Bulk TiO₂ and the P25 were quickly dispersed in the culture media when prepared for the cell exposure, which may have improved cell surface contact and subsequent uptake. Anatase and rutile were individually aggregated quickly in the culture media. This possibly reduced the particle surface area available, and consequently decreases the cell surface contact. This is in agreement with Ryabchikova et al. (2010) who showed that TiO₂ crystal structure alters particle membrane interactions which can either increase or decrease particle internalisation. They hypothesised that the NPs may alter the function of the plasma membrane with simple mechanical binding to membrane macromolecules. Furthermore, the behaviour of TiO₂ in culture media might be related

to increased ionic strength from high salt concentrations in the culture media, which may lead to aggregation of a portion of NPs (Koeneman et al., 2010).

5.4.3 Effects of the different forms of TiO₂ on Caco-2 cells electrolytes

Despite the fact that cell culture media is enriched with magnesium and calcium, exposure of Caco-2 cells to different forms of TiO₂ caused an increase in cellular magnesium and calcium concentrations and a decrease in potassium concentration in cells compared with the controls (Table 5.2). The increase in Mg^{2+} level was not observed after bulk TiO₂ exposure; this elevation in Mg²⁺ level was just seen after exposure to different forms of TiO₂ NPs suggesting a specific material effect on Mg homeostasis. Increase the accumulation of cellular Mg²⁺ requires proper distributions of the ions across the cell membrane and any changes in the membrane potential especially in the polarized epithelia like intestine as well as changes in pyridine nucleotide levels will promote cellular Mg²⁺ accumulation (Romani, 2011). This suggests that the exposure to TiO₂ may result in changes in the membrane potential. Additionally, the decrease of the potassium concentration in cells exposed to different forms of TiO_2 could be related to this change in the membrane permeability associated with the mechanical interaction of TiO₂ and membrane macromolecules (Ryabchikova et al., 2010). This potassium reduction might also be happening through the activation of single stretch-activated potassium channels which are more permeable than other ion channels in the intestinal epithelium (Chang and Loretz, 1992). The same situation as Mg^{2+} was seen with the concentration of Ca^{2+} in the cells, where the bulk TiO₂ did not cause a marked increase in Ca^{2+} concentration compared to that caused by different forms of TiO₂ NPs. Scherbart et al. (2011) suggests that calcium influxes after exposure to different particle size of TiO₂ are nanoparticle type specific. Differences between bulk and nano TiO₂ could be based on the cell sensitivity to specific material effects on the calcium homeostasis. This intracellular calcium homeostasis might be related to the

over expression or under expression of calcium-binding proteins (Fisichella et al., 2012). The highest concentration of Ca^{2+} was observed with P25 exposure (Table 5.2), which was associated with the altered structural organisation of the cell surface villi (Fig. 5.9). The same alteration in villi structure has been noticed by Koeneman et al. (2010) who relate this effect to the increase in free cellular calcium concentration.

5.4.4 How does the Ti from different forms of TiO_2 can be taken up by Caco-2 cells?

Caco-2 cells were pre-incubated for 1 h with either 120 IU ml⁻¹ nystatin (an endocytosis inhibitor) or 100 μ mol l⁻¹ sodium orthovanadate (P-type ATPase inhibitor) prior to being dosed with 1 mg l⁻¹ of different forms of TiO₂ for 24 h. These inhibitors were applied to investigate whether TiO₂ uptake is a cholesterol-dependent endocytic mechanism or an energy-dependent process.

The nystatin exposure resulted in increased cellular Ti concentrations from all forms of TiO₂ in Caco-2 cells (Fig. 5.8). Nystatin is one of the endocytosis inhibitors which inhibit the lipid-raft mediated endocytosis including the caveolae by depleting the cholesterol from the plasma membrane (Nabi and Phuong, 2003; Ragnarsson et al., 2008). Endocytosis inhibition can reduce the transport of particles across the cell monolayer (Ragnarsson et al., 2008). Increased cellular Ti concentrations may indicate that endocytosis of TiO₂ is cholesterol independent, and uptake of TiO₂ particles could be mediated by another mode of endocytosis. Exposure of prostate PC-3M cells to TiO₂ NPs indicates that cell internalization could be via clathrin-mediated endocytosis, caveolin-mediated endocytosis and macropinocytosis (Thurn et al., 2011). Ryabchikova et al. (2010) showed clusters of TiO₂ NPs within large folds of the plasma membrane in TEM images from MDCK cells, suggesting possible uptake by macropinocytosis as well as clathrin-mediated endocytosis.

The increase in cellular Ti concentration from TiO_2 in cells incubated with nystatin (Fig. 5.8) can be attributed to the blocking of some exocytosis pathways that

are cholesterol dependent. This is because the cholesterol depletion increased NP uptake by 300% (Dombu et al., 2010) and this could be also explained by the fact that cholesterol depletion is totally blocking NP exocytosis (Dombu et al., 2010). However, the exocytosis mechanism of the NPs is still not well studied. Simon et al. (2011) suggests that TiO_2 NPs can enter the cell through different internalization pathways which interact with the golgi apparatus, and can be extruded from the cell by exocytosis via endosome or secretory vesicles.

Caco-2 cells after exposed to vanadate showed further elevation in cellular Ti concentration after exposure to all forms of TiO₂ with the exception of the rutile TiO₂ (Fig. 5.8). This could be due to inhibition of the active ATP metabolism in Caco-2 cells, which is required for transport of NPs through endocytic mechanism (Rangnarsson et al., 2008). Furthermore, inhibition of ATP-ase may result in inhibition of the GTP-ase binding protein Rac-1 to the plasma membrane that stimulates the actin filament and membrane ruffling (Ridley, 2001), that essential for the macropinocytosis mechanism (Grimmer et al., 2002). Moreover, the vanadate may interrupt TiO₂ efflux by inhibiting Rho GTPases which play an important role in exocytosis through the organization and dynamics of the actin cytoskeleton (Cobbold et al., 2002). Further investigation is required to fully characterise the exact endocytic/exocytic pathways that are interfered with through vanadate application.

5.4.5 Effects of nystatin and vanadate on Caco-2 cells electrolytes

Caco-2 cells pre-incubated with nystatin and then exposed to different forms of TiO_2 showed an increase in cellular potassium and magnesium concentrations in comparison to the no drug TiO_2 controls (Table 5.2). As mentioned above, nystatin is a cholesterol depletion drug and changing the cholesterol content in the plasma membrane may cause an alteration in the membrane permeability (Rodal et al., 1999). Since the effect of TiO_2 in the cells is to reduce the level of K⁺ concentration, the elevated levels of K⁺ in the

cells after exposed to nystatin suggest that this elevation is solely caused by the nystatine incubation. For the Mg^{2+} , the elevation after exposure to nystatin was more than that after TiO₂ exposure without drugs. This suggests that the elevated level of Mg^{2+} may interfere with the effects of nystatin by altering the cell electrolytes homeostasis after cholesterol depletion.

Cells incubated with vanadate showed a decrease in both potassium and magnesium cellular concentrations compared to the no drug TiO₂ controls (Table 5.2), but an elevation was noticed in cellular Ca²⁺ concentration. This could be explained by the structural differences in the protein and crystallite. E2 was designated for the low energy phosphoenzyme which includes the Na⁺, K⁺, ATPase, Mg²⁺ ATPase and Ca²⁺ ATPase (Rice et al., 2001). Vanadate is a P-type ATPase inhibitor which inhibits both the Na⁺, K⁺, ATPase (Cantly et al., 1978) and Mg²⁺ ATPase (Rice et al., 2001) by making the transition state for phospholyration-dephospholyration mimic the state in E2. Ca²⁺-ATPase has different crystallite structure that required EGTA (Stokes and Lacapere, 1994), this may lead to more stability of these crystals which are not affected by vanadate inhibitor.

5.4.6 Presence of particles within the Caco-2 cells

The different forms of TiO₂ showed different effects on the microvilli growth on the cell surface (Fig. 5.9) indicated by a lower density of the microvilli after exposure to both P25 and rutile. The same observation was reported by Koeneman et al. (2010) who suggested that the increase in free cellular Ca²⁺ concentration was the reason for the alteration of the microvilli organisation. The current study showed the same increase in Ca²⁺ concentration after exposure to P25 (Table 5.2). The alteration in the microvilli organisation may be related to the effects of Ca²⁺ elevation on the villin, a Ca²⁺ regulated actin binding protein that is associated with the brush border microvilli cytoskeleton (Bretscher and Weber, 1980). This effect is caused by the P25, a mixture

of anatase and rutile, rather than the individual crystal forms indicating that P25 is more toxic than the other forms of TiO_2 .

The images from the SEM study showed clearly the penetration of the Ti from different forms of TiO₂ which was observed underneath the cells and in contact with the cell membrane (Fig. 5.10) and sometimes near to the nucleus (Fig. 5.11). This suggests that the NPs have the ability to enter the cells by an active transport mode which is mediated by a form of endocytic pathways. The same suggestion was made by Brun et al. (2011) after observing the appearance of TiO₂ NPs by TEM images in Caco-2 cells after 24 h exposure. They reported anecdotally, particles mainly grouped together inside the cells into heterogeneous clusters of 1-2 μ m, and hypothesised that the NPs internalization occurred through macropinocytosis.

The intracellular trafficking of NPs towards the nucleus might be happening by microtubules mostly through dynein-dependant of the endosomes processing (Panariti et al., 2012). Additionally, the shape and charge may play an important role in localization of the NPs near to the nucleus. Rode shape NPs are likely to be moved towards the nucleus via microtubules (Xu et al., 2008). Furthermore, positively charged NPs are more easily approach the nucleus (Yue et al., 2011).

Conclusions

This study shows that Ti uptake in Caco-2 cells from different forms of TiO_2 was steady, saturable and time dependent over the period of 24 h exposure; this indicates that an active absorption mechanism has occurred. The highest uptake rate was observed with the bulk TiO_2 which was about 2 fold more than that indicated by the *in vivo* study on rats (Jani et al., 1994). The direct contact of Ti from different forms of TiO_2 with the cell membrane, results in localisation of Ti inside the cell both close to the membrane and also in the cytoplasm. This suggests that the mechanism of TiO_2

uptake involves a number of different active endocytic pathways. Increased cellular Ti concentration after exposure to both nystatin and vanadate may also be related to interference with the endocytic processes. The nano form of TiO_2 showed an alteration in cell electrolytes homeostasis. P25 produced the highest increase in cell Ca²⁺ concentration which altered the microvilli organisation on the cell surface. This raises the concern that P25 toxic effects should be considered as a potential hazard to human health.

Chapter 6

General Discussion

Currently there is a very limited data on the uptake of TiO_2 NPs via intestine, the mechanisms of their uptake, and the material-type effects on Ti uptake (differences between bulk and nano). This thesis aimed to give a better understanding of the TiO_2 NPs uptake by the intestine and the possible mechanisms involved, leading to a better understanding of the hazard and risk assessment of TiO_2 NPs intake by food.

This work reports regional differences in Ti accumulation from both bulk and nano forms of TiO₂ by vertebrate intestine (Chapter 3), and demonstrates the uptake of Ti from TiO₂ particles across the gut of rainbow trout using the isolated gut perfusion technique (Chapter 3). The main findings are that Ti from TiO₂ particles (bulk or nano) are mainly absorbed in the mid and hind regions of trout intestine, with the Ti accumulating in the mucosa rather the underlying muscularis. This uptake of the Ti from TiO_2 leads us to further investigate on the possible mechanism uptake. The effects of nystatin which inhibits the uptake of Ti from TiO₂, suggest at least part of the mucosal influx step is by endocytosis for both bulk and nano forms of TiO_2 (Chapter 4). However, the effect of solute transport inhibitors also suggests a dissolved Ti component involved in Ti export from the tissue to the blood for bulk and nano TiO₂ (Chapter 4). There was also a material-type effect where Ti from NPs crossed the gut faster than Ti from the bulk powder, and uptake from the nano form was more sensitive to nystatin treatment as well as changes in the gas mixture in the media. A novel effect of the partial pressure of carbon dioxide (pCO_2) on Ti uptake is also proposed, with several possible mechanisms by which CO₂ might modulate Ti uptake from particles (Chapter 3).

In human intestinal epithelial cell line (Caco-2 cells), the Ti from all different forms of TiO_2 were uptake by the cells and the accumulation of Ti was effected by the material-type and crystal structures (Chapter 5). Bulk TiO_2 shows the greater uptake rates relative to the other forms of TiO_2 . Cells incubated with nystatin or vanadate

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demonstrate further increases in cellular Ti concentration from all forms of the TiO_2 (Chapter 5) which indicate a different situation from the intestinal tissue (Chapter 4).

Histopathology and ultrastructural (TEM and SEM) studies of the intestine for both cell and tissue provided an indicator of the uptake of Ti from different forms of TiO₂ (Chapter 3-5).

6.1 Correlation steps between the in vivo and the in vitro methods

In vivo studies are complicated by a number of considerations during dietary exposure, such as the interaction of NPs with the food components, the transit time from the mouth through the GIT, and the varying conditions between each part of the gut (pH, enzymes, gas partial pressures and composition of the digesta). All these factors could be reasonably controlled to investigate Ti uptake using *in vitro* methods.

In this study, different *in vitro* methods with different organisms were used to evaluate them with the *in vivo* studies. The studies try to mimic the conditions in real organisms and offer a direct contact between particles and intestinal epithelium. The isolated perfused intestine method is closest to the *in vivo* condition by having four layers of the intestine tissue (from the mucosa to the serosa), as well as a good viability criteria of the tissue outside the living organisms. Human intestinal Caco-2 cell is another *in vitro* method used to improve our knowledge on the uptake of NPs across a monolayer intestinal cell.

The results from both *in vitro* methods in this study gives some similarities to the results of *in vivo* studies, from the uptake of bulk TiO_2 by rat intestine (Jani et al., 1994), TiO_2 NP uptake by intestine in mice (Wang et al., 2007) and the Ti accumulation in the gut from TiO_2 NPs exposure in rainbow trout fish (Ramsden et al., 2009). However, some differences were also noticed between the *in vitro* and *in vivo* methods especially for the electrolyte concentration. Both *in vitro* methods showed alteration in gut electrolyte homeostasis which is different from what was reported by *in vivo* studies. For example, Ramsden et al. (2009) did not report any effects of TiO_2 exposure on tissue electrolytes (Na⁺, K⁺ and Ca²⁺).

6.2 Uptake and accumulation of Ti from TiO_2 exposure by the perfused intestine and Caco-2 cells

Both *in vitro* methods showed that intestinal Ti accumulation from the bulk TiO₂ was greater than the other TiO₂ nano forms. For the perfused intestine, the bulk material accumulation in the tissue was greater by 7 and 2 folds for the mid and hind intestine respectively compared to the TiO₂ NP (P25) over 4 h exposure (Chapter 3, Fig. 3.7). This was expected; the results from the serosal perfusate indicated a slower efflux of the Ti from the bulk TiO₂ across the gut to the blood side relative to the P25 resulting in Ti accumulated in the tissue. The same situation was shown in Caco-2 cells; bulk TiO₂ showed greater accumulation by 2, 6 and 8 folds compared to P25, anatase and rutile, respectively over 24 h exposure (Chapter 5, Fig. 5.7). This matches the result from the hind intestine in the perfusion experiments, in which the accumulation of bulk TiO₂ was 2 fold greater than that of the P25.

The accumulation of the Ti metal from TiO_2 exposure in gut epithelium is explained by apical uptake of the Ti in the particulate form rather than the dissolved Ti metal from different types of TiO_2 . This is supported by low proportion of the surfacebound total Ti metal on the gut mucosa that represented by rapid solution dipping experiment (Chapter 4, Fig. 4.3), and also by the measurable appearance of apparent dissolved total Ti metal from TiO_2 shown by the dialysis experiment with the normal physiological saline (mucosal solution). Furthermore, the particles in both methods were observed inside the epithelial cells using the TEM for perfused intestine and SEM for the Caco-2 cells. For the Caco-2, the SEM work was supported by using the x-ray energy dispersive spectrophotometry (EDS) measurements which detected the Ti particles in/on the cells and within the nucleus of the cells (Chapter 5, Fig. 5.9, 5.10 and 5.11). The EDS measurement with the monolayer cells is much easier to detect the particles inside the epithelial cells rather than the folded tissue layers of the intestine.

For the gut perfusion tissue, the EDS measurements can detect Ti metal in large aggregates of TiO₂ on the surface of tissues (e.g., energy dispersive x-ray on top of rodent skin, Adachi et al., 2010), but these techniques have limitations for NPs inside cells (see review, Handy et al., 2012). In traditional fixed TEM specimens, there is a limitation of EDS represented by the energy beam which can only penetrate about 2-300 nm into the tissue, detection limit of EDS compared to the concentrations of the TiO₂ used here (1 mg l⁻¹, even if it was all absorbed into the cells) would still be far below the mass detection limit against the high carbon background in the specimen (e.g., even using 10% TiO₂, the carbon peaks in Adachi et al. 2010 are off the scale compared to the much smaller, variable, Ti peaks in the spectra). A further limitation of EDS is the minimum area or cluster size to reliably detect chemical composition of the particles relative to the size of the beam (spot size between 1-50 µm). Given these technical limitation we did not observe large enough aggregates in the cells, close enough to the surface, to make reliable Ti composition measurements by EDS. However, TEM instruments especially adapted with new field emission guns for EDS are now becoming available that can achieve beams in the nm range (as used by Michel-Jeanjean et al., 2012 on skin explants), albeit with the risk that the high brightness needed may damage the specimen. Nonetheless, like previous studies with electron microscopy images of putative TiO₂ NPs inside cells without EDS measurements (e.g., Galloway et al., 2010; Powell et al., 2010) inferences may be made from particle morphology.

In the TEM images of the gut perfusion tissue, some electron dense particulate materials was detected inside the epithelial cells and sometimes located within a vesicle. These particles represent a similar morphology to the particles in the TiO_2 NPs or bulk (Chapter 3, Fig. 3.9 and 3.10), the distinct shape and size of these particles with their absence from the controls (no indication of the same particles appearance) suggesting they were a TiO_2 particles from the exposures. This observation suggests that TiO_2 NPs or bulk could enter the cells through a form of endocytosis.

6.3 Possible mechanism uptake of Ti from TiO₂ across the intestinal epithelium

The mechanism uptake of metals and metal oxide NPs is likely by endocytosis and an active uptake. The main mechanisms are clathrin-mediated and caveolaemediated endocytosis in addition to phagocytosis and macropinocytosis (Fig. 6.1) (Iversen et al., 2011; Fröhlich and Roblegg, 2012). The endocytosis of NPs is depending on the size, shape and the charge of the NPs, as well as on the cell type (Thurn et al., 2007). The general diameter of the clathrin-coated vesicles is nearly to 120 nm, while the caveosomes range between 50 and 80 nm (Johannes et al., 2002). Thus, it is likely for the NPs to be uptake by one of these endocytic mechanisms. Macropinosomes are the largest one with a range of diameter between 500 to 2000 nm (Johannes et al., 2002) which is possible for the bulk particles and the big aggregations of the NPs to be uptake by this mechanism.

The different crystal forms with so many shapes and sizes of the TiO_2 NPs possibly affecting on the endocytic pathways of its uptake which is not yet to be fully understandable. Also, the differences between the monolayer cells and the gut tissue are playing an important role in the particles uptake.

The present study indicated that the mechanism uptake of TiO_2 NPs across the epithelial cell of intestine involved one of the possible endocytic mechanisms

mentioned above (Fig. 6.1) after the sensitive responses towards the nystatin and vanadate inhibitors. Nystatin is a cholesterol depletion component via the lipid rafts and the TiO_2 NPs here acts differently towards the nystatin in the perfused intestine and Caco-2 cells. For the perfused intestine, the mechanism uptake of TiO_2 NPs could be through caveolae disturbance by cholesterol reduction and this could increase the uptake of NPs by the clathrin pathway. Thurn et al. (2011) demonstrated that uptake of TiO_2 NPs in prostate cancer PC-3M cells involved the endocytic pathways via clathrin-mediated endocytosis, caveolin-mediated endocytosis, and macropinocytosis.

In Caco-2 cells, cholesterol could be implied in the exocytosis of NPs, and for this reason the accumulation of the TiO_2 was increased after the cells were exposed to the nystatin. The cholesterol via lipid rafts may play an important role in the regulation of the exocytosis (Salaun et al., 2004; Chintigary et al., 2006). In addition, caveolaemediated endocytosis has been suggested to play a role in the uptake of NPs (Thurn et al., 2011), but if this uptake is reduced by cholesterol-depleting agents and these agents may also affect the other endocytic mechanisms (Doherty et al., 2009). These results were agreed with the study by Dombu et al. (2010) which showed clearly the implied of the cholesterol in NP exocytosis.

Exposure to nystatin and vanadate completely abolished the uptake of the bulk TiO_2 by the perfused intestine. This made a suggestion of the possible phagocytosis involvement in the endocytic uptake of the big particles of TiO_2 by phagocytic cells like macrophages (Fig. 6.1) or by the macropinocytosis within the early endosome.



Figure 6.1 The possible endocytic mechanisms uptake of TiO_2 NPs (purple dots) or bulk TiO_2 (light blue dots) showing the process of their taken up by endocytosis within the early endosome or the phagosome and both of them are then combine with the lysosome. Four active mechanisms uptake via transcellular pathway are shown in the diagram. Phagocytosis is depends on the actin filaments as a receptor mediated and largely district to M cells of Peyer's patches. This process is used for the uptake of large particles. Clatherin-mediated endocytosis is a dynamin-dependent mechanism which is also a receptor mediated uptake process. Macropinocytosis is an active, actin-dependent process used to internalize the fluid surrounding the cell containing particles then taken up altogether and located inside the cell. Caveolin-mediated endocytosis is associated with protein caveolin which is essential for the endocytosis uptake by pinch off from the membrane of the intestinal epithelium.

6.4 Material-type effects on the electrolytes of the intestinal tissue and cell

The exposure to bulk and nano TiO₂ caused some effects on intestinal electrolytes homeostasis (Mg^{2+} , Ca^{2+} and K^+). Tissue and cell electrolytes showed difference effects between the perfused intestine and Caco-2 cells methods. For the isolated perfused intestine, the bulk TiO₂ caused an alteration in tissue electrolytes homeostasis (Chapter 3, Table 3.8). This was different for the Caco-2 cells, mainly all TiO₂ nano forms was responsible for the alteration of the cell electrolytes homeostasis (Chapter 5, Table 5.2). These effects should be studied carefully which may indicate that Ti metal may interfere with the endogenous electrolytes homeostasis, especially for the Ca^{2+} which raise the concern of the ability of Ti metal to interfere with the calcium homeostasis. Further investigations on the precise mechanisms of TiO₂ induced changes in electrolyte homeostasis are required.

6.5 Hazard screening of the TiO₂ uptake by food to the GIT tract

Little is known about the potential risk of nanotechnology applications to the human body, especially within the gastrointestinal tract which is a target system for food applications. The bulk and nano TiO₂ are metal oxides that are used, or proposed, for many applications, especially in the food sector. There is also a concern that environmental releases of TiO₂ NPs may enter the food chain and reach higher trophic levels including humans. The concentrations of TiO₂ NPs in surface water have been reported to reach 0.02 μ g l⁻¹, while the concentration in the sewage treatment water was 4 μ g l⁻¹ (Gottschalk et al., 2009). For the intake of TiO₂ by food, Powell et al. (2010) estimate ingestion of 5 mg TiO₂/person/day with an unknown part of it in nanoform. Total dietary intake only of nano-TiO₂ is estimated to be 2.5 mg/individual/day (Lomer et al., 2000). Recently, Weir et al. (2012) reported that daily intake of TiO₂ by food is between 1-3 mg/Kg body weight/day, with 36% of the particles in nanoscale form. Consequently, this daily intake raises the concern about the expected of chronic effects rather than acute toxic effects on humans (Weir et al., 2012).

The present thesis highlights the possibility of hazard via the food, as relevant TiO_2 NPs uptake within the intestinal epithelium has been demonstrated. The results indicated the ability of the Ti metal from TiO_2 exposure to across the intestine, with some material-type effects on the Ti metal uptake and accumulation from TiO_2 exposure (both bulk and nano treatment); and evidence of Ti appearance inside the intestinal epithelial cells. This raises the hazard potential of the Ti accumulation from nano TiO_2 exposure which should be reconsidered in risk assessments, in addition to that based only on the bulk material. The Ti metal uptake rates from TiO_2 exposure in present study are identical to other metals like Cu (Handy et al., 2000) but it is lower by 10 fold than the Hg uptake in the perfused intestine (Holye and Handy, 2005). Thus the absorption hazard of TiO_2 is similar to Cu but potentially less toxic than Hg.

Furthermore, the effects of CO_2 in the uptake of the TiO_2 NPs through intestine also need to be considered for the risk assessment purposes especially for the human health and the pCO_2 effect on Ti uptake can be explained by acid-base toxicity in the intestine. Carbonated fizzy drinks which contain large amount of CO_2 , when combined with food intake, may accelerate the uptake of NPs through the intestine. In addition, the role of the CO_2 in elevating the uptake rate of Ti metal from TiO_2 NPs represents an important effect of these NPs in real ecosystems, associated to the partial pressure of the gasses in external medium which raise the concern of the environmental hazard from TiO_2 exposure. All the results in this study indicate an active absorption mechanism for TiO_2 in the intestinal epithelium, and this should be considered in risk assessments and dietary exposure to the TiO_2 NPs for both the ecosystems and human health. However, further investigations on the TiO_2 effects on food-related risk assessments and NPs hazards are advisable.

6.6 Future work

A number of areas of work require further investigation, and some suggestions are indicated below.

- Further investigation is required on the mechanism uptake of TiO₂ NPs by using other inhibitors of endocytic pathways. Chlorpromazine is mainly used as an inhibitor of clathrin-mediated endocytosis and known to function as a calmodulin antagonist. To further elucidate if the uptake of NPs is by a macropinocytosis pathway which is an actin-dependent mechanism, amiloride and calmodulin is often used to repress this process. Caveolae-mediated endocytosis can be inhibited by using tyrosine kinase inhibitors like genistein. More investigations on the precise endocytic/exocytotic mechanisms in conjunction with the sub cellular distribution are in demand.
- Gut rat perfusion is an ideal model to achieve our findings from the gut fish perfusion which represent a comparative study between lower vertebrates and mammals. Different concentrations of the TiO₂ (higher and lower than the 1 mg Γ^{-1}) are recommended, to learn more about the cellular uptake and intracellular transport of TiO₂ NPs across the gastrointestinal tract. Other approaches are needed to investigate more deeply about the material-type effects (particle size and/or crystal structure) on the uptake of TiO₂ by the intestinal epithelium. More knowledge on this aspect is important to determine the bioavailability of the ingested Ti from both bulk TiO₂ and the nano forms.
- To obtain more information about the uptake of TiO_2 NPs across epithelium tissue and subsequent biological effects, using the gill perfusion *in vitro* model in order to interpret and aid our data from the gut perfusion. This will enable further understanding on the uptake and behaviour of these particles by studying different parameters like the presence of organic matter such as dissolved

organic matter (DOM) which could affect NP aggregation. The salinity and hardness of the water may also alter the bioavailability and therefore the uptake of NPs. For example, the additions of Ca^{2+} to water will increase the hardness of water and that may affect the uptake and transport of NPs by influencing the aggregation of NPs, an effect which could be removed by using the Ca^{2+} chelator (+EGTA). Furthermore, the addition of NaCl will alter the ionic strength and may cause particle aggregation (reduced bioavailability). The influence of altering the water pH should be also addressed carefully, for at pH 5 the chemistry of the water and the aggregation of NPs may be different compared with pH 7.

• To assess our information about the uptake of TiO₂ NPs in either Caco-2 cells or gut tissue, it will be very useful to study molecular gene expression changes and identify the role of individual genes involved in the uptake of TiO₂ NPs. The genes that may interpret the TiO₂ uptake by gastrointestinal tract are dynamin 1, caveolin-1 and PTRF (cavin). This will provide a better understanding on the uptake mechanisms for NPs across the gut epithelium.

Research in all these aspects will increase our understanding on TiO₂ toxicity.

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Appendix I

Published Papers

REVIEW PAPER

Effects of manufactured nanomaterials on fishes: a target organ and body systems physiology approach

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(Received 11 October 2010, Accepted 6 July 2011)

Manufactured nanomaterials (NM) are already used in consumer products and exposure modelling predicts releases of ng to low $\mu g l^{-1}$ levels of NMs into surface waters. The exposure of aquatic ecosystems, and therefore fishes, to manufactured NMs is inevitable. This review uses a physiological approach to describe the known effects of NMs on the body systems of fishes and to identify the internal target organs, as well as outline aspects of colloid chemistry relevant to fish biology. The acute toxicity data, suggest that the lethal concentration for many NMs is in the mg l^{-1} range, and a number of sublethal effects have been reported at concentrations from c. 100 μ g to 1 mg l⁻¹. Exposure to NMs in the water column can cause respiratory toxicity involving altered ventilation, mucus secretion and gill pathology. This may not lead, however, to overt haematological disturbances in the short term. The internal target organs include the liver, spleen and haematopoietic system, kidney, gut and brain; with toxic effects involving oxidative stress, ionoregulatory disturbances and organ pathologies. Some pathology appears to be novel for NMs, such as vascular injury in the brain of rainbow trout Oncorhynchus mykiss with carbon nanotubes. A lack of analytical methods, however, has prevented the reporting of NM concentrations in fish tissues, and the precise uptake mechanisms across the gill or gut are yet to be elucidated. The few dietary exposure studies conducted show no effects on growth or food intake at 10-100 mg kg⁻¹ inclusions of NMs in the diet of O. mykiss, but there are biochemical disturbances. Early life stages are sensitive to NMs with reports of lethal toxicity and developmental defects. There are many data gaps, however, including how water quality alters physiological responses, effects on immunity and chronic exposure data at environmentally relevant concentrations. Overall, the data so far suggest that the manufactured NMs are not as toxic as some traditional chemicals (e.g. some dissolved metals) and the innovative, responsible, development of nanotechnology should continue, with potential benefits for aquaculture, fisheries and fish health diagnostics. © 2011 The Authors

Journal of Fish Biology © 2011 The Fisheries Society of the British Isles

Key words: behaviour; blood cells; cardiovascular system; gastrointestinal tract; nanoparticles; nervous system.

Nanotoxicology, 2012; Early Online, 1–20 © 2012 Informa UK, Ltd. ISSN: 1743-5390 print/1743-5404 online DOI: 10.3109/17435390.2012.735268

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Uptake of titanium from TiO₂ nanoparticle exposure in the isolated perfused intestine of rainbow trout: nystatin, vanadate and novel CO₂-sensitive components

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Abstract

Nanoparticle (NP) uptake across the gut is poorly understood. *In vitro* gut sac preparations and isolated perfused intestines were used to investigate the absorption mechanism(s). Exposure of whole gut sacs to 1 mg/l TiO₂ NPs for 4 h caused total Ti metal concentrations to increase in the intestine, with 80% or more of the Ti in the mucosa. Perfused intestines showed a saturable time-dependent accumulation of total Ti, which increased when the CO₂ in the gas mixture was lowered to 0.5%. Adding cyanide did not stop Ti uptake, and 100 µmol/l vanadate (ATPase inhibitor) caused a 2.8-fold reduction in the net uptake rate of Ti for TiO₂ NP exposure. Luminal additions of nystatin (endocytosis inhibitor), blocked the uptake of Ti from both bulk and TiO₂ NP treatments. The data demonstrate Ti uptake across the intestine from TiO₂ NP exposures, involving CO₂-dependent and nystatin-sensitive mechanisms.

Keywords: gut epithelium, uptake mechanism, endocytosis, carbon dioxide, dissolution

Appendix II

Poster Presentations



AG), crystal structure was 25 % rutile

and the mean primary particle size was

Before the experiments all fish were starved for 48 h. Rainbow trout,

The intestine was serosally perfused

(Figure 1) and gassed with a standard

were analysed for the lactate dehydrogenase enzyme (LDH) activity

perfused intestine

and used as viability criteria for the

Uptake of TiO₂ Nanoparticles Across the Isolated Perfusate Intestine of Rainbow Trout (*Oncorhvnchus mvkiss*)

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0.12 Hind Intestine -⁶ ¹ TiO₂ NPs, "Aeroxide" P25 (DeGussa 0.10 (umol and 75 % anatase (Federici et al., 2007) 0.08 T102 22.8 ± 0.6 nm (mean ± S.E.M., n = 169). 0.06 lissue 0.04 Oncorhynchus mykiss, weighting 239 ± 9 g (mean \pm S.E.M., n = 35) were used. 0.02 0 00 Control TIO 2N P Bulk TiO2 [TiO 2] 1 m g l⁻¹ gas mixture (95 % O₂ :5 % CO₂) or low CO_2 (99.5 % O₂ :0.5 % CO₂) for up to 4 h.

Figure 3. The TiO₂ accumulation in the mid and hind gut after 4 h perfusions with a concentration of 1 mg H⁺ for TiO₂ NP or bulk TiO₂ gassed with standard gas mix. Data are means \pm S.E.M. * Statistically significant difference (Kruskal-Valia) etc., P < 0.05 from the control value. # Bulk statistically significant differences from TiO₂ NP value. Letter (a) indicates statistically significant differences between mid and hird intestine.

* # a

Mucosal [TiO ₂]	Net TiO (nn	₂ flux, <i>J</i> _{net,TiO2} nol g⁻¹ h⁻¹)	Net water flux, J _{net,H2O} (mi g ⁻¹ h ⁻¹)				
(Initial rate	Initial rate Overall rate		Overall rate			
Standard gas mix Control	0.38 ± 0.26	0.43 ± 0.07	-6.22 ± 0.85	-7.50 ± 1.18			
Standard gas mix TiO ₂ NP	2.58 ± 0.55 *	3.97 ± 1.13 *	-17.56 ± 4.00 *	-18.01 ± 4.16 *			
Standard gas mix Bulk TiO ₂	1.64 ± 0.78 #	1.42 ± 0.54 #	-26.28 ± 2.52 *#	-28.54 ± 3.19 *#			
Low CO ₂ Control	0.85 ± 0.24	0.1 ± 0.05	-3.42 ± 1.71	-3.97 ± 0.97			
Low CO ₂ TiO ₂ NP	35.26 ± 31.5	5.45 ± 2.66	-17.13 ± 2.37 *	-22.3 ± 6.41			

Table 1. Net flux rates for TiO₂ and water across the perfused trout intestine. Values are means \pm S E M (Low CO₂, n = 6, standard gas mix, n = 7. Buk TiO₂, n = 6) expressed per gram dry mass of intestine. Negative values indicate a net loss from the serosal solution, initial and overall rates data ware calculated from curulative perfusals data at 10 min and 4. In respectively². Statistically significant difference (Kruskal-Wallis test and *t*-lest, P < 0.05) from the control value. # Bulk statistically significant difference from TiO₂ NP value.

- This study shows that the isolated perfused intestine preparation works with
- nanoparticles, and that the viability criteria remain good with these materials.
- This study also demonstrates that NPs have the ability to cross the gut of fish into
- the serosal side.
- There is a particle-size effect as the nanoform is taken up faster than the bulk powder
- TiO₂ NP absorption is time-dependent, saturable and the uptake mechanism is
- affected by the gas mixture used.

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Introduction

* The mechanisms uptake of nanoparticles (NPs) across the epithelia of fish are unknown. Handy et al. (2008) argues that apical entry of intact NPs into the epithelial cells of fish (gills, gut) is unlikely to occur via the traditional ion transport pathways (e.g., by diffusion through epithelial ion channels), because the NPs are simply far too large, but uptake into the cells by endocytosis remains possible.

✤ The aim of current study was to determine which part of the gut was involved in the uptake of TiO₂ NPs by using isolated whole gut sacs (Hoyle and Handy 1995), and then measure the uptake rate using the isolated perfused intestine (Handy et al., 2000), including pharmacological vestigations of the mechanism(s) involved.

Materials and Methods

★ 1 mg 1⁻¹ TiO₂ NP was used and prepared from a stock solution of 10 g 1⁻¹ TiO₂ NPs, "Aeroxide" P25 (DeGussa AG), crystal structure was 25 % rutile and 75 % anatase, the mean primary particle size was 22.8 \pm 0.6 nm (mean \pm S.E.M., n = 169).

S.E.M., n = 169). \diamond For the whole gut sac experiment, the gut was sutured closed at the posterior end, and the lumen filled with 1 mg l⁻¹ TiO₂ (NP, Bulk, or no added TiO₂ control as appropriate) in physiological saline. \diamond The intestine was serosally perfused and gassed with standerd gas mixture (95 % O₂ : 5 % O₂ or 10 w CO₂ (99.5 % O₂ : 0.5 % O₂) for up to 4 h. \diamond 10 mmol l⁻¹ cyanide (KCN) added to both the perfusate and mucosal solutions to determine if the TiO₂ NP uptake had an energy-dependent

♦100 µmol l⁻¹ sodium orthovanadate (Na₃Vo₄), an ATPase transporter inhibitor, added to the serosal solution of the perfusate. • In separate experiments, 120 IU ml⁻¹ nystatin (endocytosis inhibitor), was

idded to the muc osal solution only



Fig. 1. The effect of exposure the everted perfusate intestine to 1 mg l⁻¹ of TiO₂ added in the mucosal solution on cumulative [Ti] content of the serosal perfusate (nmol, bars) and cumulative serosal perfusate flow (ml, diamonds). Graphs are representative examples of individual perfusions from animono, on the territory of the second period of the second period of the territory of the second period of the territory of the second period peri 137 and 273.9 g, respectively.

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Fig. 2. The Ti metal from TiO₂ accumulation in the mid and hind gut after 4 h perfusions with a concentration of 1 mg l^{-1} for TiO₂ NP or bulk TiO₂ added to the mucosal solution gassed with standard gas mix or low CO₂, showed also the effects of drugs exposure (vanadate and nystatin) on the accumulation of the TiO₂ in the tissue. Data are means \pm S.E.M (n = 6-7). * Statistically significant difference from the control (Kruskal-Wallis, P < 0.05), # statistically significant difference from the bulk TiO₂ (Kruskal-Wallis, P < 0.05). Letter (a) indicates statistically significant difference between mid and hind intestine (t-test, P < 0.05).



Fig. 3. TEM images of rainbow trout gut epithelium cells (A) control, (B) 1 mg l⁻¹ bulk TiO₂, (C) 1 mg l⁻¹ TiO₂ NP, (D) 1 mg l⁻¹ TiO₂ NP gassed with 0.5 % CO₂ showed TiO₂ nanoparticles (C) uptake (NP) and bulk (Bk), mitochondria (Mt) noticed damaging in the exposure images, degeneration of the villi (Dg). Scale bar = $2 \mu m$

Table 1. Total [Ti] of gut tissue segments following the exposure to 1 mg 1-1 TiO₂ in the gut

[Ti] µmol g ¹ dry mass											
Treatments	Oesopha gus	Stomach	Pyloric caecae	Mid intestine	Hind intestine						
Control	<0.003*	0.003 ± 0.001*	<0.003*	0.004 ± 0.001 ^b	0.006 ± 0.004*						
Bulk TiO ₂	0.033 ± 0.009 ^{ab,*}	0.029 ± 0.007*.*	0.016 ± 0.003*.*	0.062 ± 0.018 ^{b,*}	0.041 ± 0.011 ^{ab,*}						
TiO ₂ NP	0.027 ± 0.009%*	0.017 ± 0.003*	0.015 ± 0.003*.*	0.027 ± 0.005**	0.030 ± 0.006 ^{x,*}						
We have a second s											

tissue. Whole gut sacs were filled with physiological saline (control), 1 mg l^{-1} TiO₂ NP and 1 mg l^{-1} Bulk TiO₂ at laboratory temperature (18 °C) for 4h.

* Statistically significant difference from the control value within columns (Kruskal-Wallis or ANOVA, P < 0.05). Difference letters (within rows) indicate a statistically significant difference between regions of the gut (Kruskal-Wallis or ANOVA, P < 0.05).

Conclusions

- The isolated perfused intestine and the gut sac preparations works with NPs.
 TiO₂ NP were mainly absorbed in the mid and hind regions of trout intestine, and mostly concentrated in the mucosa layer.
- TiO₂ NP absorption is time-dependent, saturable and the uptake mechanism is affected by the gas mixture used and faster uptake than the bulk powder.
- * The mechanism was found by testing inhibitors of the main uptake pathways in the gut which indicated a nystatin-sensitive pathway that may involve endocytosis in the intestine

Acknowledgments

- Al-Jubory were financially supported by the Ministry of Higher Education and Scientific Research via the Embassy of the Republic of Iraq.
- Many thanks for Peter Bond for his technical help in electron microscopy studies.

Uptake and Retention of TiO₂ Nanoparticles in Caco-2 Cells; Implications for Human Health

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Introduction

- Titanium dioxide powder is used extensively in a wide variety of commercial products. Estimates indicate that people in the western world ingest around 5 - 50 mg / day. in vivo researchs has suggested that uptake of nano $\rm TiO_2$ occurs, although little is known about the rates and mechanisms of this process.
- The aims of this study were to measure the uptake rates of ${\rm TiO}_2$ bulk powder and $\dot{\phi}$ nanopowder of differing crystal types, assess the toxicity, and, elucidate the uptake mechanisms in Caco-2 cells.

Materials and Methods



ar K⁺, Ca²⁺, Mg²⁺ and Na²⁺ in nmol [Metal] mg⁻¹ protein following exp Table 1. Tot



- ◆ 1 mg l⁻¹ TiO₂ NP containing media was prepared for each of the TiO₂ crystal types (Anatase, Rutile, and a mixture of both "P25") at the beginning of the experiment. These were prepared from a stock solution These were prepared from a stock solution of 500 mg 1⁻¹.The primary particle size measurement of 103.2 ± 16 nm, 22.8 ± 0.6 nm, 16.4 ± 2.4 nm, and 30.8 ± 2.5 for bulk, P25, anatase and rutile, respectively. A bulk particle control of 1mg l⁻¹ was used for comparative results.
 Further experiments investigated the effect
- of a range of inhibitors on particle uptake mechanism (vanadate, nystatin, chloropromazine and genistein).
- Cell health and morphology was visually inspected throughout the experiments. Membrane integrity was assessed using LDH leakage (low leakage indicates an uncompromised membrane). Cells were elementally analyzed using
- ICP-OES in order to assess the effect of TiO_2 exposure on intracellular electrolytes. cells were analysed using SEM, TEM, X-ray microanalysis, and block face analysis techniques in order to confirm that ICP detected TiO2 was intracellular in nature



the

 $\label{eq:Table 2.} Total intracellular \ K^*, Ca^{2*}, Mg^{2*} \ and Na^{2*} \ in nmol \ [Metal] \ mg^{-1} \ protein \ after exposed to \ drugs with \ 1mg \ l^1 \ TiO_2 \ polymorphs \ for \ 24 \ hrs.$

mg I ⁴		Electrolytes (ninol ing " protein)														
	K+				Ca2+				Mg2+			Na 2+				
	No drug	Chloro pr omazin e	Gashte	Andori de HCL	No drug	Chlon pr omazin e	Gealuse in	Anderi de HCL	No drug	Chioro pr omazin e	Gashte	Amiliori de HCL	No drug	Chloro promaz ine	Gasha is	An de HC
Conrol	694.2 m 165	562# 162#	60.5± 313	109.8 # 2737	15.5a 1.6	12.5± 1.3#	48.3 # 8.1~	104.1# 17.57	40.2 m 1.1	37.3 ± 1	38+a 1.6	840 m 5.07	6007.a 359	1364.a 160#	3858# 396-	+11 +57
Della	650.2 ± 31.5	555.3 # 403#	634.6 ± 35.8	977.1 ± 1997	28.5# 5.1	12.5# 1.6#	37.# 2.9~	173.9# 69.8†	40.7 ± 0.6	38.5 * 29	426 ± 1.7	704 # 7.5†	5671 ± 474	1265± 136#	4058.m 257~	370 900
P25	625.4 * 19.7	136.4 # 222#	686.5 * 36.5	771.7 + 1207	32.1 * 7.3	11.7# 1.5#	38.2 * 7.7-	86.4 * 41.17	41.5 = 2	362+ 17	43.7 ± 2.7	684 * 3.77	5659 * 365	1100+ 73#	4080.# 342~	333 802
Assuse	638.5 a 25	58.3 # 33.2#	653.7 4 33.6	795 A 1997	26.94 4.9	12.24 0.8#	40.5 A 2.4-	70.1 A 16.27	39.5 ¥ 0.9	38.6 # 24	424 A 0.8	7144 1227	5528 4 477	1010# 50#	4176a 208-	335 117
Rutle	600.5 # 343	550.9 a 185#	66.4 a 193	827.1 # 1217	17.9a 1.4	10.1# 1.2#	36.1 # 7.1~	68.6 m 11.17	37.7 m 2.1	38.3 #	439 a 0.2	799 # 8.37	4251 a 337	1041a 454	4874a 843	306

Microscopy





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Results

- TiO₂ polymorph uptake occurred in a time dependent saturable manner. Rates of uptake seem to be influenced by crystal type. Bulk and P25 NPs exhibited the most prolonged uptake and subsequent retention reaching saturation at 8 hrs (Fig. 2).
- Cells incubated with either nystatin (cholesterol binding agent) or vanadate (P-type ATPase inhibitor) ÷ caused an increase in retention relative to the no drug controls
- Incubating with genistein (tyrosine kinase inhibitor) and chloropromazine (Clathrin mediated endocytosis inhibitor) caused a decrease in TiO₂ retention.
- $\rm TiO_2$ polymorph exposure caused changes in total intracellular metal concentrations relative to non dosed cells. All Ti polymorphs caused an increase in $\dot{\mathbf{v}}$ intracellular magnesium and calcium and a decrease in intracellular potassium relative to the control.

Conclusions

- This study demonstrates that TiO_2 NPs are taken up in a time dependent saturable process.
- Uptake rates are affected by particle size and crystal structure. 4
- NPs have the ability to cross the apical membrane of Caco-2 cells and this is mediated by energy \sim ÷.
- dependent mechanisms. Drug incubation studies indicate predominant method of TiO₂ uptake is mediated by tyrosine kinase / phosphatas regulation implicating calveolae mediated mechanisms as a major uptake nathway
- TEM images show intracellular vesicles occupied by nanoparticle aggregates supporting the idea that an endocytic mechanism¹ is responsible for uptake (Image E)
- 1 mg l^{-1} doses of TiO₂ NPs are sub lethal however electrolyte levels are altered which may contribute to pathologie

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Acknowledgment

Constantinos Gitrowski and Aliaa R. Al- Jubory are equal were financially supported by the Ministry of Higher and Scientific Research via the Embassy of the Republic of