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Understanding the complexity of sediment residence time in rivers: Application of Fallout Radionuclides (FRNs)

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1	Understanding the Complexity of					
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3	Application of Fallout Radionuclides					
4	(FRNs)					
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18 Abstract

19 Riverine sediments play an important role in the healthy functioning of river ecosystems as they provide 20 nutrients and a connectivity signal throughout the catchment sediment cascade. However, excess sediment 21 supply to rivers can have several detrimental impacts on water quality, availability and ecology. The application 22 of catchment management practices requires a comprehensive understanding of both spatial and temporal 23 sediment dynamics to tackle point and diffuse river pollution. While the sources, pathways and fate of eroded 24 sediment in river systems have been widely studied, temporal dynamics have received less attention, mainly 25 due to the complexity of the processes and the lack of methods available to assess these dynamics. This 26 contribution reviews the application of Fallout Radionuclides (FRNs) as sediment residence time tracers. We 27 explore their suitability as sediment chronometers in rivers, the relevance of sediment residence time to 28 sediment budgeting, and discuss the current models that have been employed to determine sediment residence 29 time in river systems. Our review also identifies the challenges, opportunities and the future research needs for a comprehensive application of FRNs to evaluate sediment residence time. In evaluating approaches to sediment 30 31 residence time, we have summarised several pitfalls requiring consideration and identified avenues for further research. For instance, attention should be given to sorption behaviour when using ⁷Be and ¹³⁷Cs as residence 32 33 time tracers in rivers under changing environmental conditions; particle size effects; activity concentration 34 dilution by mixing of newly tagged ⁷Be sediment with ⁷Be-poor sediment from older or different sources; source 35 controls on ⁷Be delivery to rivers; and the influence of direct fallout into stream channels. Finally, further 36 research is needed to assess the influence of environmental and anthropogenic factors on sediment residence 37 time such as land use, topography, flow regimes, soil type, soil erosion measures and climate change, all of which 38 have important implications from a catchment management perspective.

39 Keywords

40 Sediments; Rivers; Residence Time; Fallout Radionuclides, Contaminants; Sediment Storage.

41 **CRediT author statement**

E. Munoz-Arcos: Conceptualisation, Formal Analysis, Investigation, Resources, Data Curation, Writing – Original
Draft, Visualisation, Funding acquisition; G. E. Millward: Conceptualisation, Writing – Review and Editing,
Supervision; C. C. Clason: Writing – Review and Editing, Supervision; C. Bravo-Linares: Writing – Review and
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administration, Funding acquisition.

47 1 Introduction.

48 Riverine sediments, typically particles < 2 mm in diameter, play an important role in the healthy functioning of 49 river ecosystems since they provide nutrients and contribute to habitat quality (Kemp et al. 2011; Jones et al. 50 2012; Wharton et al. 2017). However, human activities such as reservoir and dam construction, land use 51 changes, deforestation, mining activities and urbanisation have altered sediment fluxes in river systems resulting 52 in enhanced, or reduced, sedimentation and contamination (Walling and Fang 2003; Owens et al. 2005; Syvitski 53 and Kettner 2011; Wohl 2015). Excessive sediment supply into rivers has detrimental impacts on water quality 54 (e.g. turbidity) which causes sedimentation in river channels, reservoirs and estuaries affecting aquatic habitats 55 such as salmonids spawning gravels (Kemp et al. 2011). In addition, silts and clays are geochemically active and 56 consequently are responsible for the transport of contaminants, including trace metals, phosphorus, Polycyclic 57 Aromatic Hydrocarbons (PAHs), pesticides, radionuclides and Polychlorinated Biphenyls (PCBs) which have high 58 sorptive affinity for fine-grained sediment particles (Owens et al. 2005; Wohl 2015; Alewell et al. 2017; Rügner 59 et al. 2019; Owens 2020).

60 Fluxes of sediments in fluvial systems transport materials between hillslopes and floodplains, riparian zones, the 61 active channel, and the hyporheic and groundwater zone providing a connectivity signal within the river 62 landscape. However, it is well known that rivers are efficient in trapping sediments, and consequently only a 63 fraction of the eroded sediment may be transported to the basin outlet (Walling 1983; Fryirs 2013). Therefore, 64 sedimentation in river corridors drives channel morphology, and enhanced sedimentation in river channels can 65 have several implications including bed fining, channel narrowing allowing vegetation encroachment, altered bedform type or dimensions, bed aggradation, altered channel planform and enhanced floodplain 66 sedimentation (Wohl 2015). Furthermore, sediments can have adverse impacts on channel-beds, such as the 67 68 filling of interstitial spaces between coarser streambed sediment (also known as streambed colmation) which 69 provides habitats for aquatic insects and fish eggs, and facilitates hyporheic exchange (Kemp et al. 2011; Jones 70 et al. 2012; Wharton et al. 2017). Therefore, understanding when and how streambed colmation takes place 71 and the quantity of sediment storage, and the dwell time, are fundamental to the assessment of the 72 environmental impacts of upstream erosion and sediment pulses into rivers. Hence, quantifying the timescales 73 of sediment transfer throughout a river system is critical for understanding both river basin sediment dynamics 74 and the fate of adsorbed contaminants. This timescale is represented by the period that sediments spend within river basins, which can encompass days to months (Matisoff et al. 2005; Le Gall et al. 2017), decades (Wallbrink
et al. 2002) and millennia or longer (Hoffmann 2015).

77 1.1 Definitions

78 Finding a consensus on the meaning of sediment residence times in rivers is problematic since definitions are 79 often dependent on the methodology. For example, Whipp & Ehlers (2019) define sediment residence time as 80 the "time sediments remain within the catchment as part of a population of minerals that could be dated in a 81 random sample, assuming no long-term sediment storage in the catchment". Alternatively, Skalak & Pizzuto 82 (2010) define sediment residence time as "the time required to remove a mass of sediment equal to the total 83 volume in storage". Other authors make a distinction between terms such as sediment transit time and sediment 84 residence time. For example, Gellis et al. (2019) define sediment transit time as "the time it takes for sediment 85 to travel from a starting point in the watershed to an endpoint where sediment leaves the area of interest", 86 whereas residence time refers to "the mean time sediment particles spend in a storage reservoir". Gellis et al. 87 (2019) also noted that these terms can depend on the spatial scale used. Thus, transit time can encompass the 88 time from when sediment is released to when it enters the river channel (e.g. Slattery et al. 2002) or the time 89 that sediment spends in all storage reservoirs i.e. hillslopes, channel-bed and floodplains (e.g. Hoffmann 2015). 90 On the other hand, residence time can span sediments deposited at a specific storage unit i.e. channel-bed (e.g. 91 Skalak and Pizzuto 2010) or also include the suspended fraction (Evrard et al. 2010; Smith et al. 2014). In addition, 92 Matisoff et al. (2005) used the term sediment age to refer to the time between particles receiving their 93 radioactive tags to when they are sampled in the river as suspended, bed-stored, or estuarine sediment.

94 For consistency throughout this text with regards to defining sediment temporal dynamics the terminology 95 outlined in Figure 1 will be used. Once sediment enters a stream channel it may be transported either through, 96 or stored temporarily in, different riverine units (e.g. river bed, bars, lateral deposits and short-term floodplain 97 deposits). The time this sediment spends in these temporary units is referred to here as its storage time. 98 Sediments from these storage units can be released during stormflow events contributing to the suspended load 99 which also comprises sediments from upstream sources. Therefore, the time encompassing from upstream 100 sediment release to the time spent in riverine temporary storage and in transport is referred to here as the 101 residence or transit time. Finally, the time that a sediment particle has been resident in a long-term storage unit 102 towards the outlet of a river is given by its age (e.g. long-term floodplain deposits, lacustrine and estuarine

deposits). Therefore, sediment age spans the time that a particle has been in temporary storage, transport and
 deposition in the long-term storage unit. Although it is recognised that particles may be reworked during storage
 in these long-term deposition units, they are rarely moved to upstream catchment areas (e.g. during a tidal

106 surge).



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Figure 1. Relationships between the different concepts used here to describe the temporal dynamics of sediment storage and transport in rivers. The storage time refers to the time that sediment spends in a short-term deposition unit, residence/transit time refers to the time that sediment spends in temporary storage and in transport throughout the river system, and sediment age refers to the time that sediment has remained stored in long-term deposition units.

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113 1.2 Methods to determine sediment residence time

Several methods have been used to assess sediment temporal dynamics in different catchment compartments (Voepel et al. 2013; Hoffmann 2015; Sutfin and Wohl 2019; Carretier et al. 2020). Here we focus our discussion on riverine processes that encompass temporary storage and transport i.e. sediment residence/transit time. To date, a variety of methods have been developed to determine sediment residence time. For example, Wang Jin et al. (2015) estimated fine-grained sediment residence time using the post-earthquake rate of sediment export (measured using daily suspended sediment discharges) triggered by landslide sediment mobilised by the Wenchuan earthquake, China. Furthermore, Voepel et al., (2013) determined a bed elevation time series using 121 sonar transducers and LiDAR to evaluate empirical sediment residence times. Radionuclides offer the possibility 122 of determining both sediment age and residence times. For instance, Uranium-series isotopes have been used 123 to document the time-dependence of weathering ages (Dosseto et al. 2014; Suresh et al. 2014) and ¹⁴C dating 124 has been applied to estimate the mean sediment age of floodplain sediments (Sutfin and Wohl 2019). Shortlived radionuclides (e.g. ⁷Be, ²¹⁰Pb, ¹³⁷Cs and ²³⁴Th) have also been used to assess sediment travel distances, 125 sediment age and sediment residence time in a variety of landscapes (Olsen et al. 1986; Dominik et al. 1987; 126 127 Wieland et al. 1991; Vogler et al. 1996; Bonniwell et al. 1999; Feng et al. 1999; Ciffroy et al. 2003; Forster et al. 2009). In lakes, Wieland et al. (1991) modelled sediment residence times using ⁷Be and ²¹⁰Pb fluxes in Lake 128 129 Zurich, Switzerland, and Vogler et al. (1996) estimated sediment residence times in Lake Constance for the total radionuclide inventories of ²³⁴Th and ⁷Be. In estuaries, Olsen et al. (1986) assessed water column removal rates 130 131 and residence time of ⁷Be in the James River Estuary, USA, and Ciffroy et al. (2003) determined sediment 132 residence time of suspended particles using ⁷Be budgets in the turbidity maximum zone of the Loire estuary, 133 France. The application of these radionuclides as sediment residence time tracers in lakes, coastal and estuarine zones has been discussed elsewhere (Dominik et al. 1989; Steinmann et al. 1999; Feng et al. 1999; Baskaran 134 2001; Baskaran and Swarzenski 2007; Forster et al. 2009; Saari et al. 2010). For extensive reviews on the use of 135 136 radionuclides in these environments the reader is directed to works from Du et al. (2011); Kaste et al. (2002); 137 Kaste & Baskaran (2011); Waples et al. (2006).

138 *1.3 Rationale and objectives of this review*

139 Increasing attention given to the temporal dynamics of fine-grained sediment in river systems (Du et al. 2011; 140 Walling 2013; Matisoff 2014; Collins et al. 2020) raises the need for a revision of the use of medium and short-141 lived radionuclides as chronometers, the methods applied, the identification of potential limitations, and 142 implications from a catchment management perspective. If the temporal dynamics of suspended and channel-143 bed sediments are better understood, then the timeframe when catchment management practices become 144 effective can be better constrained, or otherwise, to justify the difficulty to implement mitigation plans over 145 short timescales. This is a key aspect to consider when developing mitigation actions to solve river basin point 146 and diffuse pollution, and the subsequent ecological consequences that fine sediment storage and remobilisation can have on rivers. In this context, this review primarily addresses the use of short-lived 147 148 radionuclides to assess sediment residence time in rivers to: 1) assess the existing literature regarding sediment residence time with a focus on the use of Fallout Radionuclide (FRNs) tracers in river systems; 2) provide an assessment of the main sediment residence time models applied to date; 3) discuss the assumptions and challenges of these different methods; and 4) identify future research needs for a comprehensive evaluation of sediment residence time in river systems.

153 2 River sediment budgeting and its relevance to channel storage and

residence time.

155 Sediment dynamics in river systems involve complex processes, and their quantitative assessment faces many uncertainties. River sediment budgeting has become increasingly used to overcome this issue because it 156 provides an understanding of the sediment mobilisation, transport, storage and yield (Walling and Collins 2008). 157 158 River sediment budgets can be understood, in simplistic terms, as the mass balance between the sediment 159 sources, deposition areas and outputs. One of the most significant findings from budgetary studies is the 160 importance, and magnitude, of sediment storage in rivers. From the total amount of sediment produced (i.e. 161 eroded) in upland surfaces (i.e. sources) only a fraction makes its way to the basin outlet (i.e. the sediment yield). 162 This discrepancy has been named the 'sediment delivery problem' by Walling (1983) and it has been argued that various sediment storage mechanisms operating within a catchment may explain this discrepancy (Trimble 1983; 163 164 Walling 1983; Fryirs 2013). Since then, an expanding body of research has been carried out to explain and 165 disentangle the mechanisms, pathways and fates of eroded sediment within a river basin, however, the 166 temporal dynamics of these processes have received less attention due in part to the complexity of the problem 167 and the lack of available methods.

An issue related to sediment budgeting concerns interpretation of the delivery ratio term (i.e. the ratio of the inputs to the output) (Walling 1983; Parsons 2012). For example, substantial variability at different temporal scales (e.g. between stormflow events, seasons or years) has been found in sediment delivery ratios within a catchment, the so-called 'temporal lumping' as described by Walling (1983). Therefore, it is necessary to accompany channel sediment budgeting with an improved understanding and evaluation of the sediment residence times within different storage units to avoid temporal biases and uncertainties in the interpretation of the sediment delivery ratio.

175 In many catchments, sediments spend significant time stored in riverine compartments, and delivery is therefore 176 controlled by storage and sporadic remobilisations over various timescales. These storage units can be defined 177 as transient, short-lived landforms, such as bars, lateral deposits and the streambed, that are frequently reworked during stormflow events where they play a key role in the (dis)connectivity of the catchment sediment 178 179 cascades (Fryirs 2013). Thus, river channel sediment budgeting becomes a key tool in the assessment of 180 sediment reworking/exchange magnitudes/quantities in rivers subjected to episodically driven sediment 181 remobilisation and deposition for a comprehensive understanding of the sediment dynamics in these temporary 182 storage units.

183 3 Suitability of medium and short-lived radionuclides as sediment

184

residence time tracers.

Medium and short-lived radionuclides (i.e. ²¹⁰Pb, ¹³⁷Cs, ⁷Be and ²³⁴Th) have been extensively used as soil and 185 sediment tracers to assess their redistribution, deposition rates and residence time (Feng et al. 1999; Blake et 186 al. 2002; Waples et al. 2006; Mabit et al. 2008; Blake et al. 2009; Mabit et al. 2013; Taylor et al. 2013; Walling 187 188 2013; Mabit et al. 2014; Taylor et al. 2019). However, it is not the aim of this review to examine the current 189 knowledge of radionuclides as soil erosion and sediment tracers, for which the reader is directed to reviews by Mabit et al. (2014, 2013); Matisoff (2014); Matisoff and Whiting (2011); Parsons and Foster (2011); Taylor et al. 190 (2013); and Walling (2013). Moreover, applications of ²³⁴Th as a tracer of sediment dynamics in freshwater 191 systems (Waples et al. 2006) have mostly concerned lakes (Dominik et al. 1989; Vogler et al. 1996; Waples et al. 192 2004). Some studies have determined ²³⁴Th in river sediments and water despite the typically low activity 193 194 concentrations (Morris et al. 1994; Waples et al. 2003) but no sediment residence time studies using this 195 radionuclide in river systems have been found. For this reason, we focus our discussion on Fallout Radionuclides 196 (FRNs).

¹³⁷Cs ($t_{1/2}$ = 30.17 years) is an anthropogenic radionuclide that was released to the stratosphere as a result of atmospheric thermonuclear weapon testing in the 1950s – 1980s, in addition there have been releases during nuclear accidents such as the 1986 Chernobyl and the 2011 Fukushima disasters. Before subsequent deposition on the Earth's surface, ¹³⁷Cs circulated globally and was washed out by precipitation patterns (Ritchie and McHenry 1990). ²¹⁰Pb ($t_{1/2}$ = 22.2 years) is of geogenic origin and is a natural decay product within the ²³⁸U decay series derived from the decay of the inert gas ²²²Rn ($t_{1/2} = 3.8$ days) which derives from its parent ²²⁶Ra ($t_{1/2} = 1622$ years). The ²¹⁰Pb generated in situ by decay of ²²⁶Ra is termed supported ²¹⁰Pb and is in equilibrium with ²²⁶Ra. However, ²²²Rn diffuses to the atmosphere and undergoes a series of short-lived decays to ²¹⁰Pb which may adsorb to aerosols and is delivered to the landscape by wet and dry fallout. Fallout ²¹⁰Pb is termed unsupported, or excess ²¹⁰Pb (²¹⁰Pb_{ex}). ⁷Be ($t_{1/2} = 53.3$ days) is a cosmogenic radionuclide produced by cosmic ray spallation of nitrogen and oxygen in the stratosphere and troposphere and it is delivered to the earth's surface through wet and dry deposition.

An advantage of using ¹³⁷Cs, ²¹⁰Pbex and ⁷Be as sediment chronometers, is that the contrasting half-lives that 209 210 these radionuclides possess can be used to model sediment residence time from days to decades in different catchment compartments (Wallbrink et al. 2002; Matisoff et al. 2005; Le Cloarec et al. 2007; Gellis et al. 2019). 211 Additionally, naturally and continuously produced ⁷Be and ²¹⁰Pb radioisotopes can be used at many sites in the 212 world widening their application (Wallbrink et al. 2002; Matisoff et al. 2005; Le Cloarec et al. 2007; Jweda et al. 213 2008; Evrard et al. 2010; Smith et al. 2014). The basis for the use of these radionuclides as tracers is their ability 214 215 to rapidly and strongly adsorb onto soil and sediment particles following both wet and dry deposition (Welp and Brümmer 1999; Du et al. 2011; Matisoff and Whiting 2011; Taylor et al. 2012; Matisoff 2014; Singleton et al. 216 2017; Ryken et al. 2018) with reported partition coefficients, K_d, of the order 10⁵ (Olsen et al. 1986; Hawley et 217 218 al. 1986; Dominik et al. 1987; Van Hoof and Andren 1989; You et al. 1989; Steinmann et al. 1999; Kaste et al. 2002; Jweda et al. 2008). However, sorption behaviour of ¹³⁷Cs and ⁷Be have been questioned, especially in 219 changing environments through the catchment sediment cascade which can alter sediment residence time 220 221 estimations (Parsons and Foster 2011; Taylor et al. 2012, 2013; Ryken et al. 2018). For example, partitioning of ⁷Be and ¹³⁷Cs have been reported to be strongly influenced by pH and salinity (You et al. 1989; Kaste et al. 2002; 222 Giannakopoulou et al. 2007; Hong et al. 2011; Kaste and Baskaran 2011; Taylor et al. 2013), therefore, its 223 224 application in estuarine environments as a sediment particle tracer is challenging and requires case-by-case 225 validation.

²²⁶ 4 Models to evaluate sediment residence time in river systems.

227 Models have been developed and tested to evaluate sediment residence time in rivers within a variety of 228 landscapes, catchment sizes and regions across the world (see Table 1 and Figure 2). Application of radionuclide 229 mass balances to assess sediment residence times began in late 1980s using data from the Rhone catchment, 230 Switzerland (Dominik et al. 1987). However, it was not until late 1990s that other authors employed different 231 methodologies to determine/model sediment residence times. For example, Wallbrink et al. (1998) determined 232 the proportional contribution from three potential sources: cultivated and uncultivated land and subsoil 233 material from gullies and channel banks. Then, applying a mixing model with various concentrations of ²¹⁰Pbex 234 corrected as a function of radioactive decay in the channel, they found that the mean residence time of fine-235 grained material was 10 ± 5 years (Table 1). However, evidence that sediment residence times in the 236 Murrumbidgee river, Australia could be of the order of weeks to months was noticed from changes in the mean ¹³⁷Cs activity concentrations of suspended sediments between flood and low-flow conditions, and the presence 237 of ⁷Be activity in sediments from flood water. Bonniwell et al. (1999) used ⁷Be/²¹⁰Pb and ²¹⁰Pb/¹³⁷Cs ratios to 238 239 assess the fraction of new sediment in suspension, residence times and transport distances. In their study, the basis for the use of ⁷Be/²¹⁰Pb ratios was stablished because it corrected for the relative sorption and enrichment 240 241 effects resulting from variations in grain size and particulate matter composition, and by using ²¹⁰Pb/¹³⁷Cs ratios they accounted for variations in mineralogy and source area activity concentrations. Residence times in 242 sediments from the channel of River Gold Fork, USA, ranged from 1.6 to 103 days (Table 1) from the upper to 243 244 the lower part of the catchment. In addition, Wallbrink et al. (2002) determined residence times using ²¹⁰Pbex activity concentrations in river bed sediments from the Brisbane and Logan rivers, Australia. They assumed that 245 increases in ²¹⁰Pbex/¹³⁷Cs ratios in sediments occurred because of primarily direct input of fresh flux of ²¹⁰Pbex to 246 247 the sediments in the river channel. This was supported because no such higher ratios were measured at potential soil erosion sources within the catchment. Using the initial inventory of ²¹⁰Pbex within the mobile layer of river 248 249 sediment derived from the catchment erosion, ²¹⁰Pbex inventories at the time of sample collection and the inventory of ²¹⁰Pbex which occurs when depositional flux decay within the sediment profile reach equilibrium, 250 251 they calculated residence times for deposited sediments in rivers Brisbane and Logan of 0 - 21 and 0 - 9 years 252 (Table 1), respectively. This model was also applied by Douglas et al. (2009) in the Maroochy River estuary, 253 Australia. In their study, modelled sediment residence times averaged between 1 - 28 years.

Table 1. Studies that have used FRNs to model/determine sediment residence time in the literature.

Site location	Site	Radionuclide(s)	Modelling approach	Sediment residence	Reference
Rhone Watershed,	5,220 km ²	$^{\rm 137}\rm Cs$, $^{\rm 210}\rm Pb$ and $^{\rm 7}\rm Be$	Two-box model.	800 – 1,400 y; 1 – 220 d	Dominik et al.
Murrumbidgee River, Australia.	13,500 km²	²¹⁰ Pb _{ex}	²¹⁰ Pb _{ex} source decay as a function of in-channel residence time	10 y	(1987) Wallbrink et al. (1998)
Gold Fork River, USA.	389 km²	137 Cs, 210 Pb and 7 Be	Normalised activity	1.6 – 103 d	Bonniwell et al. (1999)
Brisbane and Logan River catchments. Australia.	13,600 and 3,076 km ²	¹³⁷ Cs and ²¹⁰ Pb	Comparison of catchment soil erosion inventories to riverbed sediment inventories	0 – 21 and 0 – 9 y	Wallbrink et al. (2002)
Old Woman Creek, Weeks bay and South Slough, USA.	69.5, 24.3 and 73 km ²	^7Be and $^{210}\text{Pb}_{ex}$	⁷ Be/ ²¹⁰ Pb _{ex} ratio	50 – 300 d	Matisoff et al. (2005)
River Seine basin, France.	7 to 65,700 km²	¹³⁷ Cs, ²¹⁰ Pb and ⁷ Be	Two-box model.	115 – 307 d (river box); 4,859 – 31,192 y (soil box)	Le Cloarec et al. (2007)
Clinton River, Southeast Michigan, USA.	1,980 km²	⁷ Be and ²¹⁰ Pb _{ex}	Single-box model	0.2 – 2.1 d (⁷ Be), 0.5 – 8.6 d (²¹⁰ Pb)	Jweda et al. (2008)
Maroochy river estuary, Australia.	630 km²	¹³⁷ Cs and ²¹⁰ Pb	Comparison of catchment soil erosion inventories to river bed sediment inventories	1 – 28 y	Douglas et al. (2009)
Cointzio catchment, México.	3 to 12 km²	$^{137}\text{Cs},^{210}\text{Pb}$ and ^{7}Be	Two-box balance model.	50 – 200 d (river box); 5,000 – 23,000 γ (soil box)	Evrard et al. (2010)
		^7Be and $^{210}\text{Pb}_{ex}$	⁷ Be/ ²¹⁰ Pb _{ex} ratio	101 – 163 d	
White, West rivers and Mink Brook: USA	29 to 319 km²	⁷ Be and ²¹⁰ Pb _{ex}	Stacked reservoirs model.	4 to > 300 d	Gartner et al. (2012)
Pheasant sub- catchment, USA	12.4 km ²	^7Be and $^{210}\text{Pb}_{ex}$	⁷ Be/ ²¹⁰ Pb _{ex} ratio	40 – 319 d	Huisman and Karthikeyan (2012)
Pheasant sub- catchment, USA.	12.4 km ²	^7Be and $^{210}\text{Pb}_{ex}$	⁷ Be/ ²¹⁰ Pb _{ex} ratio	9 – 318 d	Huisman et al. (2013)
River Tamar basin, UK.	38 to 219 km²	$^{\rm 137}\rm Cs,~^{\rm 210}\rm Pb$ and $^{\rm 7}\rm Be$	Two-box model.	185 – 368 d (river box); 77,000 – 48,000 y (soil box)	(2014) (2014)
Pheasant Valley Catchment. USA.	50 km²	^7Be and $^{210}\text{Pb}_{ex}$	⁷ Be/ ²¹⁰ Pb _{ex} ratio	123 – 322 d	Lamba et al. (2015)
Loroux catchment, France.	25 km²	^7Be and $^{210}\text{Pb}_{ex}$	⁷ Be/ ²¹⁰ Pb _{ex} ratio	20 – 200 d	Le Gall et al. (2017)
Midwestern USA rivers	6.8 to 5,893 km²	²¹⁰ Pb _{ex}	Age of surface derived sediments	0 – 174 d	Gellis et al. (2017)
		^7Be and $^{210}\text{Pb}_{ex}$	⁷ Be/ ²¹⁰ Pb _{ex} ratio	61 – 282 d	, , ,
White clay creek, USA.	7.25 km ²	^7Be and $^{210}\text{Pb}_{ex}$	⁷ Be/ ²¹⁰ Pb _{ex} ratio	22 – 110 d	Karwan et al. (2018)
Orge river catchment	900 km ²	^7Be and $^{210}\text{Pb}_{ex}$	⁷ Be/ ²¹⁰ Pb _{ex} ratio	18 – 140 d	Froger et al.
Clinton River, Southeast	1,946 km²	$^{210}\text{Po}_{ex}$ and $^{210}\text{Pb}_{ex}$	Single-box model	0.3 – 3.9 d (²¹⁰ Pb), 0.9 – 13.4 d (²¹⁰ Po)	Baskaran et al.
Walnut creek	52.6 km ²	⁷ Be and ²¹⁰ Pb _{ex}	Age of surface derived	$44 - 205 d (^{7}Be),$ $1 - 58 x (^{210}Pb)$	Gellis et al.
Ducktrap River, USA.	9 km reach	⁷ Be	CIA (Constant Initial	0 to > 160 d	Fisher et al.
South River, USA	37 km reach	$^{14}\text{C}\text{, }^{137}\text{Cs}$ and $^{210}\text{Pb}_{ex}$	Reservoir theory model	1 – 60 y	Skalak and Pizzuto (2010)





258 A different approach was adopted by Skalak & Pizzuto (2010) who applied several radiometric dating methods (¹⁴C, ²¹⁰Pb and ¹³⁷Cs) to infer the distribution of ages of sediment stored within Fine-Grained Channel Margin 259 260 (FGCM) deposits of the South River, USA. By applying the reservoir theory¹ to the population of ages dated on 261 the deposits, they found a sediment residence time of 1.75 years with a very small portion of sediments with 262 storage times of the order of decades (60 years). One of the main benefits of this approach is that it provides 263 both residence time and a distribution of ages. Similarly, Fisher et al. (2010) studied sediment storage times 264 using ⁷Be coupled with a Constant Initial Activity (CIA) sediment aging model to assess transitional bedload 265 storage times in bars associated with in-channel obstructions (large wood debris and boulders). They identified 266 two dominant transport regimes with differing storage times: 1) transport-limited reaches with storage times > 100 days associated with channel obstructions, and 2) supply-limited reaches associated with steeper gradients 267 and greater stream power capable of mobilising fine-grained sediments from channel obstructions with 268 269 generally < 100 days of sediment storage. Although they addressed successfully the bed-storage time problem

¹ Reservoir theory describes the change in abundance of a substance in a reservoir in terms of its inputs and outputs through the reservoir. This reservoir can be defined as any volume enclosed by a boundary such as a lake, an ecosystem or a soil (Eriksson 1971; Mudd and Yoo 2010).

in a study of a 9 km reach of the Ducktrap River (USA), certain precautions were noted with regards to the
 application of ⁷Be when using CIA aging model:

272 1) ⁷Be activity concentration dilution by sediment depleted in ⁷Be through long-term in-channel residence
 273 times (greater decay) and/or frequent landslides or bank collapses which may supply ⁷Be-depleted
 274 sediments diminishing initial activities used to feed the CIA model, and

2) ⁷Be sediment enrichment by fresh tagging from atmospheric inputs into submerged bars at low-flow
 conditions which may overprint the inherited ⁷Be signal and thus increase initial activities of ⁷Be.

277 More recently, a novel technique to determine sediment residence time in riverbed sediment at various depths 278 was developed by Gartner et al. (2012). In their study, the channel bed is divided into a vertically stacked series 279 of reservoirs based on the assumption that residence times in stream beds would likely increase with depth, 280 yielding not a single residence time but rather a distribution of residence times varying with depth. Applying this 281 model in cores taken from the White and West Rivers (Vermont) and the Mink Brook (New Hampshire), USA, 282 they found sediment residence times of about 2 months in unregulated rivers while residence times exceed 6 months in regulated rivers. In addition, based on ⁷Be and ²¹⁰Pbex depth profiles, they suggested two mechanisms 283 284 of bed material exchange: filtration; and scour and fill. For those core profiles that showed systematic decline in 285 FRN activity a filtration mechanism was inferred, whereas scour and fill was linked to an inconsistent variation 286 in activity concentrations with depth.

As shown in Table 1, different models have been developed to assess sediment residence time in rivers, but relatively few have been applied more than once and in different settings. These models are reviewed in more depth in the following sections.

290 4.1 Single box model

The single box model described here uses FRNs to assess sediment residence times in river systems (Jweda et al. 2008; Baskaran et al. 2020). Particle residence times are obtained by means of mass balances of particulate $^{210}Pb_{ex}$ (Eq. 1) and ⁷Be (Eq. 2) to obtain sediment resuspension rates as follows (Figure 3):

294
$$\Psi_{Pb}^{d}A_{Pb}^{d} + \lambda_{Pb}A_{Rn}^{p} + I_{Pb}^{rp} + \frac{R_{Pb}A_{Pb}^{r}}{H} = O_{Pb}^{rp} + \lambda_{Pb}A_{Pb}^{p} + \frac{R_{Pb}A_{Pb}^{s}}{H}$$
 Eq. 1

295 and

296
$$\Psi_{Be}^{d}A_{Be}^{d} + I_{Be}^{rp} + \frac{R_{Be}A_{Be}^{r}}{H} = O_{Be}^{rp} + \lambda_{Be}A_{Be}^{p} + \frac{R_{Be}A_{Be}^{s}}{H}$$
 Eq. 2

Where I_{Pb}^{rp} and I_{Be}^{rp} are the input fluxes of particulate ²¹⁰Pb_{ex} and ⁷Be (dpm cm⁻³ yr⁻¹), respectively, O_{Pb}^{rp} and O_{Be}^{rp} 297 are the output fluxes of particulate 210 Pb_{ex} and 7 Be (dpm cm⁻³ yr⁻¹), respectively, H is the mean depth of the water 298 column (cm), A_{Rn}^p is the activity of ²²²Rn (dpm cm⁻³) adsorbed onto particulate matter, A_{Pb}^d and A_{Be}^d are the 299 activities of dissolved ²¹⁰Pb and ⁷Be (dpm cm⁻³), A_{Pb}^{p} and A_{Be}^{p} are the activities of particulate ²¹⁰Pb_{ex} and ⁷Be (dpm 300 cm⁻³), A_{Pb}^r and A_{Be}^r are the activities of ²¹⁰Pb_{ex} and ⁷Be in the resuspended material (dpm g⁻¹), A_{Pb}^s and A_{Be}^s are 301 the activities of 210 Pb_{ex} and 7 Be in the settling particulate matter collected in sediment traps (dpm g⁻¹), Ψ_{Pb}^{d} and 302 Ψ^d_{Be} are the first-order removal rate constants of ²¹⁰Pb_{ex} and ⁷Be (y⁻¹) from the dissolved phase onto particles, 303 and R_{Pb} and R_{Be} are sediment resuspension rates using ²¹⁰Pb_{ex} and ⁷Be (g cm⁻² yr⁻¹), respectively. 304



305

Figure 3. Diagram of the single box model illustrating the sources and sinks of particulate ²¹⁰Pb. Modified from Jweda et al.

307 (2008). The same model concept applies to ⁷Be sources and sinks but without the input from the decay of a parent

308 radionuclide.

Assuming that ²¹⁰Pb_{ex} and ⁷Be activities of the upper layer of bottom sediment are equal to those of resuspended sediment and that the production term from ²²²Rn is negligible, then sediment resuspension rates can be obtained from mass balance equations of ²¹⁰Pb_{ex} (Eq. 3) and ⁷Be (Eq. 4):

313
$$R_{Pb} = \frac{H(\Psi_{Pb}^{c} A_{Pb}^{d} - \lambda_{Pb} A_{Pb}^{p})}{A_{Pb}^{s} - A_{Pb}^{r}}$$
Eq. 3

314 and

315
$$R_{Be} = \frac{H\left(\Psi_{Be}^{c}A_{Be}^{d}-\lambda_{Be}A_{Be}^{p}\right)}{A_{Be}^{s}-A_{Be}^{r}}$$
Eq. 4

Where Ψ_{Pb}^{c} and Ψ_{Be}^{c} are the rate constants corresponding to the scavenging of dissolved ²¹⁰Pb and ⁷Be onto particles (d⁻¹), respectively. Then, particle residence times can be calculated as follows:

318
$$\tau_{Pb}^{p} = SPM \frac{H}{R_{Pb}^{r}} 365$$
 Eq. 5

319
$$au_{Be}^p = SPM \frac{H}{R_{Be}^r} 365$$
 Eq. 6

where τ_{Pb}^{p} and τ_{Be}^{p} are the residence times of particulate ²¹⁰Pb_{ex} and ⁷Be (d), respectively, SPM is the Suspended Particulate Matter (g cm⁻³), *H* is the height of the water column, and R_{Pb}^{p} and R_{Be}^{r} are the resuspension rates for ²¹⁰Pb_{ex} and ⁷Be, from Eq. 3 and 4, respectively.

323 Using the single box approach, Jweda et al. (2008) modelled sediment residence times in the Clinton river from 0.5 to 8.6 days using ²¹⁰Pb_{ex} and from 0.2 to 2.1 days using ⁷Be (Table 1). They found that particulate radionuclide 324 325 residence times were significantly lower than the dissolved radionuclide residence time. This was attributed to 326 the influence of colloidal-bound radionuclides (water samples filtered to < 0.5 μm pore size) during resuspension of bottom sediments which were accounted within the dissolved fraction. Moreover, an inverse strong 327 relationship between ²¹⁰Pb and ⁷Be log K_ds and log SPM was observed (R > 0.90) suggesting a particle-328 329 concentration effect attributed to the influence of colloidal cycling of particle-reactive species in the river. In 330 another study in the same river calculated sediment residence times ranged from 0.32 to 3.86 days using ²¹⁰Pbex and from 0.9 to 13.4 days using ²¹⁰Poex (Baskaran et al. 2020). 331

One of the advantages of this model is that it incorporates the dissolved fraction of the radionuclides (Eq. 1 and
2; Figure 3). Although it is difficult to sample and measure dissolved radionuclides, integration of this fraction

334 compensates for *in situ* particle scavenging from the available dissolved radionuclides, which in turn accounts 335 for fresh atmospheric input and radionuclide desorption from particles. Another important advantage is the 336 quantification of the sediment resuspension rate, which allows estimation of sediment reworking from the 337 storage compartment i.e. bottom sediments as in Jweda et al. (2008). This term could have potential implications 338 for elucidating the quantities of old sediment in storage that can be remobilised. This gap in knowledge has been identified as a major difficulty regarding the use of ⁷Be as tracer in sediment residence time studies (Fisher et al. 339 340 2010; Gellis et al. 2017) unless this contribution (i.e. sediment depleted in 7 Be) can be quantified within the 341 timeframes of study.

342 4.2 Two-box balance model

The two-box balance model to determine sediment residence time was first developed by Dominik et al. (1987), 343 344 later improved by Le Cloarec et al. (2007) and applied internationally (i.e. Mexico and UK) by Evrard et al. (2010) 345 and Smith et al. (2014). Here the catchment is subdivided into two boxes (Figure 4) as follows: a) the soil box (given an s subscript in the text and equations) is characterised by an area S_s with low transport velocities and 346 347 long residence times. It comprises the uppermost soil surface subjected to radionuclide fallout (Evrard et al. 348 2010), and b) the river box (given an r subscript in the text and equations) is characterised by the river area, S_r , 349 and its immediate surroundings e.g. runoff and sediment-generating areas on hillslopes connected to the river 350 network, faster exchanges and shorter residence times (Smith et al. 2014).



351

Figure 4. Conceptual diagram of the two box-model for sediment residence time estimations using FRNs. Boxes are characterised by an area S_s (soil box) and S_r (river box). Modified from Evrard et al. (2010).

354 The two-box model requires solution to four equations:

355 Firstly, the fraction of atmospheric flux, F_a , present in each box and is given by:

356 $S_s + S_r = 1$ Eq. 7

357 Then the mass balance equations for each box are required. In the soil box, the loss of material is either 358 by transport into the river box or by radioactive decay:

$$F_a S_s = I_s (k_s + \lambda)$$
 Eq. 8

360 where k_s is the rate of transfer out of the soil box, related to the residence time τ_s , with $k_s = 1 / \tau_s$; I_s is the 361 radionuclide inventory in the soil box, and λ is the constant of radioactive decay. 362 The third equation represents the addition of atmospheric inputs to the river box flux coming from the 363 soil box. Both fluxes are required to balance the output from the river and the radioactive decay:

364
$$F_a S_r + I_s k_s = I_r (k_r + \lambda)$$
Eq. 9

365 where I_r is the radionuclide inventory in the river box and k_r is the rate of output transfer from the river box.

366 Finally, the flux exported from the river box, F_r , is calculated:

$$367 F_r = I_r k_r Eq. 10$$

These four equations are written for ¹³⁷Cs, ²¹⁰Pbex and ⁷Be considering the partitioning coefficients of the 368 369 radionuclides between water and suspended matter, none of which were addressed in the former version of 370 Dominik et al. (1987). In order to solve these equations, the following assumptions based on the λ values are 371 made: 1) the radionuclide residence time in the soil compartment is expected to be of the order of decades to centuries, therefore, most of the ⁷Be will be lost by decay implying its export from the soil compartment is 372 373 negligible that is: $k_s \ll k_{Be}$ and $I_{s_{Be}}k_s \ll S_rF_{a_{Be}}$; 2) the decay rate of ²¹⁰Pb_{ex} is considered negligible compared 374 the export rate from the rapid compartment: $\lambda_{Pb} \ll k_r$. It is also assumed that the duration of sediment storage in the rapid compartment does not result in net decay of ⁷Be to levels below the limit of detection (Smith et al. 375 376 2014).

The rapid compartment area, S_r , can be determined by combining Eq. 9 and 10 based on the first assumption. Then, S_r is a function of the ratio F_r/F_a for ⁷Be and the export rate ($k_r = 1/\tau$):

379
$$S_r = \left(\frac{F_r}{F_a}\right)_{Be} \left(\frac{\lambda_{Be}}{k_r} + 1\right)$$
 Eq. 11

380 The slow box residence time, τ_s , is then computed by solving Eq. 7 to 9 for ²¹⁰Pb_{ex} with the assumption that 381 $\lambda_{Pb} \ll k_r$ and combining this with Eq. 11:

382
$$\tau_{s} = \frac{\left(\frac{1}{\lambda_{Pb}}\right) \left[1 - \left(\frac{F_{a}}{F_{r}}\right)_{Pb}\right]}{\left(\frac{R_{r}}{R_{a}}\right) (1 + \lambda_{Be} \tau_{r}) - 1}$$
Eq. 12

383 where $R_r = (F_r)_{Be}/(F_r)_{Pb}$ and $R_a = (F_a)_{Be}/(F_a)_{Pb}$.

Another relationship implicating τ_s and τ_r can be derived from the mass balance equations of ¹³⁷Cs. Given the atmospheric flux of ¹³⁷Cs is currently negligible i.e. $(F_a)_{Cs} = 0$, its inventory in soils can be determined. Based on Eq. 9, one can then determine τ_s as follows:

387
$$au_s = \frac{[M - (F_r)_{Cs} \tau_r]}{(F_r)_{Cs} (\lambda_{Cs} \tau_r + 1)}$$
 Eq. 13

388 where $M = (I_s)_{Cs} + (I_r)_{Cs}$.

The river box residence time, τ_r , may then be obtained by combining Eq. 11 and 12 and finding a solution to the equation:

391
$$a(\tau_r)^2 + b(\tau_r) + c = 0$$
 Eq. 14

393
$$a = -(F_r)_{CS} \left(\frac{R_r}{R_a}\right) \lambda_{Be} \lambda_{Pb}$$
,

394
$$b = M\left(\frac{R_r}{R_a}\right)\lambda_{Be}\lambda_{Pb} - (F_r)_{CS}\lambda_{Pb}\left(\frac{R_r}{R_a} - 1\right) - (F_r)_{CS}A\lambda_{CS}$$
 and

$$395 \qquad c = \lambda_{Pb} M\left(\frac{R_r}{R_a} - 1\right) - A(F_r)_{CS}$$

396 in which

$$397 \qquad A = 1 - \left(\frac{F_a}{F_r}\right)_{Pb}$$

398 The two-box model has been applied in various catchments with catchment surface areas ranging from 7 to 399 65,000 km² (see Table 1) and sediment residence times ranging from 4,800 to 30,321 years in the soil box and from 50 to 365 days in the river box. Le Cloarec et al. (2007) found a strong positive relationship between the 400 soil box residence time and the catchment area within the Seine basin (see Figure 5, where R = 0.98, p < 0.001), 401 402 while no relationship was found between the river box residence times and the catchment surface area. A 403 correlation analysis of residence times modelled using the two-box model in the available literature 404 demonstrates that this case is an exception, and we find no significant relationship between these two variables 405 in the literature (Figure 5). Interestingly, an inverse relationship (R < -0.6) was found in the River Tamar basin, 406 UK, between river box residence time and catchment surface area. However, only the 2007 period for the Tamar 407 was significant (p < 0.05). This finding suggests that residence times are, to some extent, independent of the 408 catchment area, and that local factors such as topography, land use and climate may play a significant role in 409 sediment release, transport and storage. Nevertheless, comparison between these studies is not straightforward 410 as methods for data collection are different and some correlations are influenced by single extreme values. In 411 this case, more studies (and data) are needed to better assess the influence of various catchment features on 412 sediment residence time.



413

Figure 5. Correlation analysis between catchment area and soil box and river box residence times for data extracted from
Evrard et al. (2010); Le Cloarec et al. (2007) and Smith et al. (2014) on the Cointzio, Seine, and Tamar river basins, respectively.
Note that every data point (n = 8 for Seine basin, n = 3 Cointzio catchment, and n = 7 for River Tamar basin) represents a subcatchment with a given area within the basin.

418

Sensitivity analysis was carried out by Le Cloarec et al. (2007) to assess the two-box model performance (Figure
6), and uncertainty simulations showed an important variation of sediment residence times for the soil and river
boxes when suspended matter concentrations (i.e. suspended load) were modified (Figure 6a, b). The effect of

422 changes in this parameter on the estimated residence times were approximately a factor of 2. Variation in the atmospheric fluxes of ⁷Be and ²¹⁰Pb (i.e. F_a) did not significantly influence soil box residence times (τ_s) (Figure 423 424 6d, f), whereas a notable variation was found in the river box residence times (τ_r) (Figure 6c, e). For example, variations in the atmospheric flux of ²¹⁰Pb (about 1.5 times) decreased residence time by a factor of 2 in the river 425 426 box. The influence of the ¹³⁷Cs catchment inventory was not as important as previous parameters when 427 modelling river box residence times (Figure 6g), but it exerts an important control when modelling soil box 428 residence times (Figure 6h). Overall, sensitivity analyses showed that estimation of suspended sediment fluxes, ¹³⁷Cs inventory, and atmospheric fluxes of ⁷Be and ²¹⁰Pb exert important controls on model outputs. 429 430 Furthermore, it was also found that ignoring the radionuclide Kd could influence residence time estimations by a factor of 2 (Le Cloarec et al. 2007) and by factors of 1 - 1.3 (Smith et al. 2014). 431





433 Figure 6. Sensitivity analysis of river and soil boxes residence times (τ_r and τ_s), illustrating the response of modelled residence times by changing parameters such as suspended matter (SM) concentration (a, b), atmospheric fluxes (F_a) of 434 435 ²¹⁰Pb (c, d) and ⁷Be (e, f), and the ¹³⁷Cs catchment inventory (g, h) in different sub-catchments of the Seine River basin. 436 Extracted and modified from Le Cloarec et al. (2007).

437 4.3 The ${}^{7}Be/{}^{210}Pb_{ex}$ ratio as an indicator of sediment age or the fraction of new sediment in suspension

Another method used to determine sediment residence time includes the evaluation of the ⁷Be/²¹⁰Pb_{ex} ratio as
an indicator of sediment age (hereafter sediment residence time as explained in section 1.1) or, alternatively,
the fraction of new sediment in suspension (Matisoff et al. 2005).

441 In this approach, sediment residence times are calculated as follows:

442
$$t = \frac{-1}{\left(\lambda_{7_{Be}} - \lambda_{210_{Pb}}\right)} \ln\left(\frac{A}{B}\right) + \frac{1}{\left(\lambda_{7_{Be}} - \lambda_{210_{Pb}}\right)} \ln\left(\frac{A_{0}}{B_{0}}\right)$$
Eq. 15

443 where $\lambda_{7_{Be}}$ and $\lambda_{210_{Pb}}$ are the decay constants of ⁷Be and ²¹⁰Pb, respectively, A and B are the activity 444 concentrations of ⁷Be and ²¹⁰Pb_{ex} in the sediment samples (e.g. suspended and/or channel bed sediments) 445 respectively, and A₀ and B₀ are the activity concentrations of ⁷Be and ²¹⁰Pb_{ex} in the source, respectively. The 446 source term may refer to activity ratios from precipitation (Matisoff et al. 2005) or sediments from overland flow 447 (Le Gall et al. 2017). The second term in Eq. 15 is a constant with a value determined by the ⁷Be/²¹⁰Pb_{ex} ratio in 448 the source.

449 Alternatively, the contribution of recently eroded particles can be calculated as follows:

450 % "new" sediment =
$$100 \times \left(\frac{A_{B}}{A_{0}/B_{0}}\right)$$
 Eq. 16

451 The ${}^{7}\text{Be}/{}^{210}\text{Pb}_{ex}$ chronometer application is carried out under several assumptions:

452 1) dry and wet fallout are included although ⁷Be and ²¹⁰Pb_{ex} are delivered to the landscape primarily during
 453 precipitation events;

- 454 2) ⁷Be and ²¹⁰Pb_{ex} are delivered to the soil in a constant proportion regardless season, latitude or proximity
 455 to the ocean and variations in the atmospheric flux is eliminated by using their ratio;
- 456 3) once deposited onto the landscape, ⁷Be and ²¹⁰Pb_{ex} are assumed to be rapidly and irreversibly absorbed
 457 to particulate matter;
- 458 4) ⁷Be and ²¹⁰Pb_{ex} are not partitioned differentially onto particulate matter; and
- since radionuclide sorption behaviour of ⁷Be and ²¹⁰Pb_{ex} is similar, differences along the flow path
 caused by particle size are eliminated by considering their ratio (including mineralogical variations).

461 The ⁷Be/²¹⁰Pb_{ex} approach to determine sediment residence time has, however, been subject to critical appraisal 462 (Walling 2013). An aspect that has been challenged is the 'source' term (i.e. the initial activity ratio) required by the method. The use of the ⁷Be/²¹⁰Pb_{ex} ratio in rainfall as the initial activity ratio was used as a constant term in 463 464 several works (Matisoff et al. 2005; Evrard et al. 2010; Huisman et al. 2013). Nevertheless, it is known that substantial variability can be found, both temporarily and spatially, in the atmospheric fluxes of ⁷Be both 465 466 between and within storm events. For example, Gourdin et al. (2014a) showed that spatial variability in activity 467 concentrations were significant within the same storm with differences up to 6-fold for ⁷Be and 4-fold for ²¹⁰Pb at different stations within the storm. Furthermore, ⁷Be/²¹⁰Pb activity ratios increased 2-fold during one storm 468 469 reflecting different controls on the fallout between these two radionuclides. Consequently, recommendations 470 of complete rainfall sampling, deposition separation of respective successive storms, spatially distributed 471 collection of rainwater, and the reduction of uncertainties related to rainfall collection were made (Gourdin et 472 al. 2014a). The assumption of constant activities derived from rainfall is thus poorly supported by the available evidence. Also Walling (2013) criticised the sediment source controls on the ⁷Be/²¹⁰Pb_{ex} ratio, since freshly 473 474 mobilised sediment will reflect the ⁷Be/²¹⁰Pb_{ex} ratio of their sources and thus their relative contributions. In this case, contributions from one source to another may change through time (i.e. within and between storm 475 476 events). For example, some sources have particulate matter with a given activity concentration of ²¹⁰Pbex as a result of an accumulated inventory and being exposed to fresh fallout of ⁷Be, therefore modifying their ratio. 477 478 One way to overcome this issue is the collection of sediments from overland flows to estimate source 479 radionuclide activity concentrations of fresh sediment inputs into the river (Gourdin et al. 2014b). This approach 480 is particularly helpful if sediment is originated primarily from catchment surface sources (Le Gall et al. 2017) but 481 the relative contribution of sources needs to be quantified.

The ⁷Be/²¹⁰Pb_{ex} ratio is the most applied method to study the temporal dynamics of sediment within river systems. Some studies have applied this method to assess the fine sediment dynamics during floods (Gourdin et al. 2014b; Le Gall et al. 2017), while others have used it to investigate the sediment dynamics and sources of sediment associated pollutants such as trace metals (Froger et al. 2018), pesticides (Gellis et al. 2017) and phosphorus (Huisman and Karthikeyan 2012; Huisman et al. 2013; Lamba et al. 2015).

487 *4.4 Age of surface derived sediments*

488 Recently, a new approach linked a sediment fingerprinting method with sediment dating in order to determine 489 the age of the surface-derived portion of sediments (Gellis et al. 2017, 2019). This method uses ⁷Be and ²¹⁰Pb_{ex} 490 for two age classes: ⁷Be up to ~ 1 year and ²¹⁰Pb_{ex} up to ~ 85 years (Gellis et al. 2019). A generalised version of 491 this model is as follows:

492 Firstly, ⁷Be activity concentrations are corrected to the percent of surface-derived sediment:

493
$${}^{7}Be_{corr} = \frac{{}^{7}Be_{target} - \left[{}^{7}Be_{s_1} \left(\frac{\% s_1}{100} \right) + \dots + {}^{7}Be_{s_n} \left(\frac{\% s_n}{100} \right) \right]}{\frac{\% surf}{100}}$$
 Eq. 17

where ⁷Be_{corr} is the estimated ⁷Be activity in the surface; ⁷Be_{target} is the measured activity in the target sample; ⁷Be_{s1} is the mean activity concentration of ⁷Be in the non-surface derived sediment source 1, if it can be measured e.g. fresh tag of ⁷Be on exposed sub-surfaces such as horizontally aligned sub-surfaces (i.e. rilled and scalded hillslopes) and/or gully areas during high rainfall and higher river flows (Wallbrink and Murray 1993; Hancock et al. 2014; Evrard et al. 2016); $%s_1$ is the % of contribution of the sediment source 1 to the target sediment sample obtained by the fingerprinting results; and %surf is the surface derived percentage obtained from the sediment source apportionment. The age of the target sediment is then computed as follows:

501
$${}^{7}Be_{age} = \frac{\ln\left(\frac{{}^{7}Be_{corr}}{{}^{7}Be_{surf}}\right)}{-\lambda_{7Be}}$$
 Eq. 18

where ${}^{7}Be_{age}$ is the age of the of the topsoil derived sediment; ${}^{7}Be_{corr}$ is the estimated surface ${}^{7}Be$ activity concentration; ${}^{7}Be_{surf}$ is the estimated ${}^{7}Be$ activity concentration of the source material (surface); and $\lambda_{7_{Be}}$ is the decay constant for ${}^{7}Be$.

505 Next, the estimated topsoil activity for ²¹⁰Pb is determined as follows:

506
$${}^{210}Pb_{ex_{corr}} = \frac{{}^{210}Pb_{ex_{target}} - \left[{}^{210}Pb_{ex_{s_1}}\left(\frac{\%{s_1}}{100}\right) + \dots + {}^{210}Pb_{ex_{s_n}}\left(\frac{\%{s_n}}{100}\right)\right]}{\frac{\%{surf}}{100}}$$
 Eq. 19

where ${}^{210}Pb_{ex_{corr}}$ is the estimated ${}^{210}Pb_{ex}$ activity in the surface; ${}^{210}Pb_{ex_{target}}$ is the measured ${}^{210}Pb$ activity in the target sample (e.g. suspended sediment); ${}^{210}Pb_{ex_{s_1}}$ is the mean activity concentration in the non-surface derived sediment source 1; ${}^{\otimes}s_1$ is the percentage of contribution of the sediment source 1; and ${}^{\otimes}surf$ is the surface derived percentage obtained from the sediment source apportionment. Finally, the age of the targetsediment is computed as follows:

512
$$^{210}Pb_{ex_{age}} = \frac{\ln\left(\frac{210Pb_{ex_{corr}}}{210Pb_{ex_{surf}}}\right)}{-\lambda_{210Pb}}$$
 Eq. 20

513 Where ${}^{210}Pb_{ex}{}_{age}$ is the age of topsoil-derived sediment; ${}^{210}Pb_{ex}{}_{corr}$ is the estimated surface ${}^{210}Pb_{ex}$ activity 514 concentration; ${}^{210}Pb_{ex}{}_{surf}$ is the weighted ${}^{210}Pb_{ex}$ activity concentration in the surface; and $\lambda_{210}{}_{Pb}$ is the decay 515 constant of ${}^{210}Pb$.

516 The age of the surface-derived sediment approach was first applied by Gellis et al. (2017) using ⁷Be in a study 517 comprising samples from 99 catchments in the Midwestern region of USA where residence times ranged from 0 518 to 174 days in bed sediments and from 0 to 84 in suspended sediments. They also found higher concentrations of pesticides (i.e. bifenthrin and DDE) in samples from the streambed with greater proportion of surface-derived 519 520 sediments and relatively young residence times (< 100 days), which gives an indication of the role that sediment 521 sources might play in the delivery of contaminated sediment and how quickly they move through the river 522 system. In a subsequent study carried out in the agricultural Walnut Creek catchment (Iowa, USA), modelled sediment residence times ranged from 44 - 205 days using ⁷Be and from 1 - 58 years using ²¹⁰Pb_{ex} (Gellis et al. 523 2019) (Table 1). In this study, sediment transport and storage were depicted in three boxes with three types of 524 ages: 1) a rapid box with less than a year (based on ⁷Be results), 2) a decadal box comprising from 10 to 100 525 years (based on $^{210}Pb_{ex}$ results) and 3) a geological box from 100 to > 1,000 years (based on the literature). 526

527 In their model, Gellis et al. (2019) defined age as the residence/transit time (see section 1.1) encompassing the 528 time from when sediments enter a channel from a surface source to when the target sample is collected. As ⁷Be 529 is used as an indicator of the age of the surface derived sediment, an assumption of surficial erosion is made, 530 and that erosion of deeper soil layers is neglected. Although this assumption was supported by a fingerprinting study, the effects of deeper erosion may be important and should be accounted for. In this regard, deeper 531 surface erosion would tend to reduce ⁷Be activity in sediments (Wallbrink and Murray 1993; Walling 2013) 532 533 which, in turn, can increase sediment residence time estimations. Gellis et al. (2017) also acknowledge that fresh input of ⁷Be on wetted channel areas of a stream channel can increase the activity of ⁷Be. It has been estimated 534 535 that direct contribution of 7 Be to large rivers could cause a 10 – 12% increase in activity concentrations (Hancock et al. 2014). Hence, if atmospheric inputs on wet areas of the channel are important, then estimated ages ofsediment might be younger.

538 5 Challenges, opportunities and future research needs.

The development of sedimentary models has contributed to an improved understanding of the temporal 539 540 dynamics of catchment sediment storage and transport, however there are still some pitfalls that require careful 541 consideration and further research. Most of these constraints are closely related to assumptions surrounding the behaviour of ⁷Be when using it as soil and sediment tracer. For instance, there is no way at present of 542 543 separating out the effects of decay and dilution of ⁷Be activity concentrations in sediments, which raises one of 544 the biggest limitations of the sediment residence/transit time methods when using FRNs (Matisoff et al. 2005; Fisher et al. 2010; Walling 2013; Gellis et al. 2019). The mixing of newly tagged ⁷Be sediment with ⁷Be-dead 545 546 sediment (e.g. sediment from channel banks and/or sediment stored in streambeds and floodplains) is thus an issue yet to be addressed. More research is needed in this regard to quantify the uncertainty, or otherwise 547 548 apportion the amount, of ⁷Be-depleted sediments that are stored in riverine compartments. Furthermore, because most of the ⁷Be activity concentration in soils is found in the top centimetre of surface soil (Blake et al. 549 550 1999; Schuller et al. 2006; Walling 2013), the catchment soil may be subjected to deeper erosion that will tend 551 to lower activity concentrations of suspended sediment which can result in overestimations of sediment 552 residence times (Gellis et al. 2019).

553 Equally important is the direct fallout of ⁷Be and ²¹⁰Pbex in wetted areas of the stream channel which may 554 increase activity concentrations of radionuclides in suspended sediments and sediments stored in the surface of 555 the bed channel. It has been reported that direct fallout of radionuclides on stream channels is negligible compared to sediment inputs from erosion of the catchment soils (Hancock et al. 2014). Although this 556 557 assumption has been supported in catchments with enhanced overland flow and surface runoff (Evrard et al. 2016; Le Gall et al. 2017), studies in forested and forest-influenced catchments have shown that channel 558 559 interception of FRNs can be important (Karwan et al. 2016, 2018). Future studies should thus acquire information regarding hydrological flow paths and hillslope connectivity to address possible problems when assessing 560 561 sediment residence times, or otherwise, quantify the proportion of radionuclide inputs into channels from

precipitation, especially at storm events. This would allow for a corrected sediment residence time thatconsiders the influence of rainfall-delivered radionuclide tags onto sediment particles.

564 ⁷Be signatures of a specific source will continue changing through time in response to radioactive decay and input of fresh fallout. It is important to recognise, therefore, that $^{7}Be/^{210}Pb_{ex}$ ratios could be expected to vary 565 566 both between storm events and seasonally, as well as from year to year, due to various controls on ⁷Be and 567 ²¹⁰Pb_{ex} fallout (Walling 2013; Gourdin et al. 2014a). Consequently, characterisation of signature of dominant 568 sources should be undertaken as a core component of any sediment residence time study. This poses important challenges when planning sampling and its related logistics, particularly due to the short half-life of the ⁷Be and 569 570 the need to encompass, as much as possible, the variability of ⁷Be in the sediment source signature (Gourdin et 571 al. 2014a).

572 Radionuclide activity concentrations may correlate with grain size (He and Walling 1996), and their effects on 573 tracer properties have been widely debated in the literature (Smith and Blake 2014; Laceby et al. 2017). 574 Therefore, selective transport of particles, in terms of their size, can strongly influence sediment residence time 575 calculations. Particle size analysis is, consequently, highly recommended in this regard, and when applicable 576 correction factors should be applied to the radionuclide activity concentrations.

577 As discussed in section 3, the sorption behaviour of ¹³⁷Cs and ⁷Be have been questioned, especially in changing 578 environments (Parsons and Foster 2011; Taylor et al. 2012, 2013; Ryken et al. 2018). Rivers are dynamic systems 579 that are subjected to rapidly changing conditions, influencing variables such as pH, redox, dissolved oxygen, temperature and conductivity. This has important implications when determining sediment residence times. For 580 example, partitioning of ⁷Be and ¹³⁷Cs between water and the particulate phase have been reported to be 581 582 strongly influenced by pH and salinity (You et al. 1989; Kaste et al. 2002; Giannakopoulou et al. 2007; Hong et 583 al. 2011; Kaste and Baskaran 2011; Taylor et al. 2013) suggesting the potential for these tracers to desorption 584 under changing environments in the wider catchment. The potential for overestimation of residence times must, 585 therefore, be considered.

586 Finally, the influence of environmental and anthropogenic factors on sediment residence time in river systems 587 are poorly understood. Only a few studies have attempted to link and quantify the relationship between 588 residence times and catchment processes under different scenarios such as changing land use (Smith et al. 2014)

- and pollution (Huisman et al. 2013; Gellis et al. 2017; Froger et al. 2018). More research is thus needed to
- 590 understand the influence of various catchment characteristics such as land use, topography, flow regimes, and
- 591 soil type on sediment release, transport, and residence time in rivers. Moreover, assessment of the impact of
- 592 soil erosion measures, catchment management practices, and climate change on sediment residence times in
- 593 river channels is still lacking but has important implications from a catchment management perspective.

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