A preventable death? A family’s perspective on an adult safeguarding review regarding an adult with traumatic brain injury

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A Preventable Death? A family’s perspective on an adult safeguarding review regarding an adult with Traumatic Brain Injury

Abstract

Purpose of this paper: To review the care management of a man with a Traumatic Brain Injury (TBI) from a family member’s perspective.

Approach: The paper provides a case history of “Tom” both prior to his TBI and after.

Findings: Tom was the subject of an adult safeguarding review in Somerset following his death in 2014. Ultimately the paper highlights the shortcomings and failures in the care Tom received by various organisations which ultimately contributed to his suicide.

Practical implications: The paper highlights the need for more effective communication between professionals managing the care of those with TBI. Furthermore, professionals need training in the need for mental capacity assessments and improved safeguarding and risk assessments with adults with TBI.

What is new in this paper: This paper provides insight into the needs of an adult with TBI from the perspective of a family member who is also a trained psychologist.
Introduction

In December 1993 my older brother “Tom” had a serious road traffic accident. This accident led him to suffer a significant traumatic brain injury (TBI) which changed his and our lives forever. Tom took his own life in June 2014. This paper will discuss Tom’s life, the impact of his head injury and provide insight into the shortcomings and failures in the care he received post-injury.

Tom prior to TBI

Tom was 10 years older than me so I didn’t really know him as a child. He moved out when I was six and had his accident when I was 12. My main memories of him are from when he was about 16 onwards. I idealised him as most little sisters idealise their big brothers. Even then he was a troubled young man with mental health difficulties, namely depression, but all I saw was a boy who loved his little sister who always wanted to spend time with her.

Tom had a different father, and the separation between our mother and his father was acrimonious. His Dad rejected him and to a certain extent, so did our Mother, leaving him feeling isolated and unloved. At secondary school Tom made lots of new friends who took away some of that sense of rejection, but they were not real friends. He was a poor working class boy in a public school environment and they used him. Before long he was getting into trouble and was ultimately expelled from school. He went on to be expelled again and then once at his third school, gave up attending.

Tom loved motorbikes and was always getting into trouble for riding too fast, or too dangerously. He was often in trouble with the police for traffic offences or other crimes involving drugs or petty theft. He was eventually assigned a social worker as result of these
activities. What he needed was love and affection and someone to listen to him, but that didn’t happen and as a result he was set on a path of self-destruction.

Tom’s upbringing meant that he often seemed depressed, although he had no formal diagnosis. He also struggled to manage his emotions and would often become frustrated and agitated. Tom also experienced a series of small, yet potentially significant, head injuries throughout his childhood that may have contributed to his risk taking behaviour, depression and difficulties with emotional regulation. At the age of four, Tom was hit by a car, and although it was at low speed and the injuries were mild, Tom did hit his head and experienced a concussion injury. Then again at the age of eight, Tom fell off his bike after riding it too fast down a hill. This resulted in him being hospitalised overnight for concussion injuries. He then went on to have a further accident with a cricket ball early on in secondary school. The significance of these multiple head injuries was missed by the clinical teams at the time, and nobody made the potential connection between these injuries, his emotional dysregulation, depression, and his increasingly risky behaviour (Sariaslan et al, 2016). Tom needed intervention at this point, and this really should have been in the form of a child psychologist who may have been able to identify the underlying issues for Tom and provide him with greater support. It may also have proven useful to us as a family by providing us with a greater understanding of Tom’s behaviour.

Tom’s Accident

On the afternoon of the 22nd December 1993, Mum and I were in town doing some last minute Christmas shopping and we bumped into Tom. We had not seen him for a while and he seemed different; positive and full of life. He had a girlfriend; they were happy and in love. He told us about a new job he had and how much he enjoyed it; he was very much
looking forward to finally turning over a new leaf. I was excited and felt I might finally get
my brother back, be able to spend more time with him and get to know him properly. Little
did I know at the time that I was right but in a way I couldn’t possibly imagine.

Later that afternoon Tom got in a car. I never saw that man again. That evening the police
informed my Mother that Tom had been involved in a serious accident and was in hospital.
They did not know what the prognosis was, but it did not look good. At that time our family
were all distraught. We just wanted Tom to wake from his coma and be with us again. In
many ways our grieving process for Tom started that day over 20 years ago.

Looking back I don’t remember much about those early weeks and months. I don’t know if
that’s because of time or if I have deliberately forgotten much of it. I know it was painful
and I felt way too young to be able to cope with what was happening to us. I remember the
main focus though. Firstly we were thinking in terms of whether he would live or die. Soon it
became clear he would live, so the focus shifted to whether he would wake from his coma.
Once he was awake, the focus shifted again to whether he would ever walk or talk again,
and what kind of life he might have. Overall we were just so pleased to have him back with
us. Unfortunately as the years progressed it became increasingly apparent that it would
have been kinder to everyone concerned, including Tom, if he had died in that accident.

Each stage of Tom’s recovery was exhausting and terrifying but filled with such sense of
hope. I can honestly say that once he started to show small signs of recovery, we really
started to believe everything was going to be alright. None of us had any idea of what was
ahead of us. This is a story that you will hear time and time again from family members of
those with brain injuries because there is simply not enough information and support in
those early months.
I do remember a couple of visits to the hospital. One was early on when Tom was still in a coma. The other was going to visit him once he had regained consciousness. He was fairly alert but couldn’t walk or talk. At that stage he couldn’t feed himself because his jaw was wired and he had hemiplegia on his dominant right side. I remember I was asked if I wanted to feed him. I cannot describe what a difficult experience that was; a 13 year old girl feeding her 23 year old brother. He sat there in silence just staring at me and I wasn’t sure he knew who I was. Overall I was very impressed with the care Tom received from the hospital in those early days. Sadly, leaving hospital and intensive rehabilitation behind marked a drastic decline in the care and support Tom received.

**Tom post TBI**

Tom left hospital in the spring of 1994 and shortly after got married to his girlfriend. This was when the reality of our situation really started to hit. Within a few months, Tom’s wife had walked out on him and we were left to pick up the pieces. This was obviously a devastating emotional blow for him but was compounded by a complete lack of support from any organisation. Tom could just about cook for himself but was not really able to function properly. However, he never received any support and it was up to us to try and guide him to keep the place clean and to look after himself. This was a difficult task as he had no sense of smell or appetite due to neurological damage, so often did not eat properly. He also had obsessive tendencies, mainly around collecting items. This meant that he would pick objects up from his walks and keep them, or refuse to throw useless items away. He was also prone to emotional outbursts, and was fiercely independent. This meant trying to tidy his house, or remove any of his newly acquired possessions (e.g. unpaired shoes or lost cuddly toys) had to be handled very carefully. For example on one occasion he returned
from town with a single black stiletto. When I suggested it should be thrown away as it
served no useful purpose, he became agitated and said to me “I have no purpose but you
aren’t throwing me away”. The shoe eventually became glued to his wall above the door in
his lounge.

Tom obviously had a history of drug abuse prior to the accident but in that first year post
injury he had remained clean and was proud of that for a short time. But it didn’t last and he
soon sought out support from a small number of his old friends, the few who were still
prepared to see him, and this inevitably led to a gradual creep back into drug use. Initially
only cannabis and alcohol but this slowly led to opium and ultimately heroin.

I had wanted desperately before the accident to know my brother better and instead he
was taken away and I was presented with a complete stranger who I was now partially
responsible for at the age of 13. As Tom’s friends had mostly abandoned him, he started to
go out with me and my friends as a way of getting some social contact. This was hard for me
as I loved him so much but he was often very embarrassing as he would be inappropriate
with my female friends as he hadn’t relearnt his social filters yet. For example one evening
when we were out, Tom grabbed the bottom of one of my close friends. She turned around
and politely asked him to let go, but was clearly agitated. Tom’s response was “well you
smiled at me, you clearly want sex”. My friend was only 15 and Tom was 25. He would also
get angry and aggressive if people didn’t understand him when he spoke, or if they
inadvertently made him feel weak and inferior. For example, on occasion a male friend of
mine held a door open for him into a pub. Tom responded by shouting “I’m not a f***ing
invalid”. This would lead to disagreements with my friends and a feeling in me of torn
loyalty. Nobody understood these behaviours and it was really only intuition on my part that
I understood this was a result of the injury to his brain and not him. Nobody ever discussed
the problems we may have along these lines and certainly offered us no support in
managing it. He hated being in a wheelchair and would really get angry about it. So he
started training himself to walk further and further each day. For months I would leave
school, walk to his flat and walk with him back and forth until he was able to navigate the
journey to the local newsagents.

Tom’s accident changed him in several profound ways. Although he had always been quite
headstrong, he had never been unusually angry or aggressive. But after the accident this
was a central part of his character. He was volatile; one minute he would be happy and
talking and the next angry or deeply depressed. He had severe memory problems but this
was often hidden behind an ability to remember very specific detailed accounts from
aspects of his life pre-injury. This often gave others the perception that his injuries were not
that severe. For example one of his crowning achievements was being able to regularly
complete the mathematical challenges on the TV show countdown. He even boasted of the
numerous times he had managed to get the answer within the time even when Carol
Vorderman could not. Yet he would regularly forget important tasks, like attending
appointments and conversations he had had with people. He also struggled with attention
and was easily fatigued. His speed of comprehension was particularly affected, especially if
he was in a group of people. He would struggle to follow a conversation and it would take
him some time to catch up afterwards. This led him to become very quiet in group
situations and only later would he announce that he had not understood what was said, or
more often, he would seize on a fairly innocuous comment and view this as evidence that
people were “out to get him” and fail to process any other parts of a conversation. For
example, on one occasion a meeting was held between the family and Headway Somerset to try and see what could be done to support Tom further. At the meeting the then Manager of Headway made several suggestions for supporting Tom and was obviously genuinely concerned. However, Tom’s take home message from the meeting was that “she wants to lock me up”. This was in reference to a comment she had made about how he may end up in prison if his drug taking did not get under control. Again this was often missed because he had maintained his intellect and seemed quite capable of expressing himself. These symptoms of TBI are often described by those who are familiar with them as hidden disabilities, and Tom did have many. But he also had very obvious physical disabilities too. Sadly at times I feel his movement problems and his shaking were put down to his drug taking rather than symptoms of his neurological damage.

During the time between 1997 and 2004 Tom found something to give his life some meaning. Tom started to attend a day centre and soon became very keen on woodwork. He spent a long time learning this new found skill, making lots of interesting and bizarre gifts for his friends and family. The day centre was also the place where he first met his long-term girlfriend.

For the next few years, things ticked over fairly normally in Tom’s life and ours. Tom’s new girlfriend was great for him. She gave his life purpose and meaning and for once something to live for. They had their many ups and downs but their companionship was unyielding. Further to that, Tom’s relationship with her also led him to come in contact with the newly formed Headway Somerset. Headway too gave Tom’s life some purpose, and was particularly important given the financial cuts at the day centre he had been attending. The team at Headway did everything they could to help Tom improve his life and were an
important source of support for me too. By this time I had formally taken on the role of Tom’s advocate, so it was my job to speak for him where necessary and to generally support him. This was not for a specific purpose, but as a general advocate to assist him in appointments and general interactions with professionals.

Although I say this was a fairly stable point, the years were still punctuated by many regular “crises”. There were drugs overdoses (deliberate or otherwise) and passing out or falling over as a result of his substance abuse, which often landed him in hospital. He would fall out with his girlfriend or his girlfriend’s family, or be threatened with eviction because he couldn’t keep his flat clean and tidy. Towards the end of 2004 it became evident to me that Tom was not doing well. I remember turning up at his flat one evening and he was sitting in his chair smoking a cigarette and I recall thinking he looked like an AIDS patient - thin, drawn and grey and I knew he needed help. His neurological recovery was going reasonably well but he had become very depressed and introverted.

Service responses to Tom’s care

In the early years following Tom’s accident, he presented on numerous occasions to the emergency department with falls as a result of his drinking. His TBI had left him with severe epilepsy and alcohol did not agree with him or his medication. We never received any contact from the hospital during this time informing us of these injuries, so he was always considered by the medical staff to not require any additional support. Even once I became Tom’s advocate I received no contact from them. He was also meant to have a social worker but we didn’t really know much about this after his initial post-hospital accommodation had been sorted. We felt utterly alone as a family. We had no support from anyone.
Tom’s declining mental health led to an assessment and appointment with the local hospital mental health services in 2005. In the initial session the psychologist discussed with Tom the principles of cognitive behavioural therapy and trying to map emotions, thoughts and behaviours. We spent some time over the next two weeks trying to piece together Tom’s triggers. When we went back for the next session it became apparent that it had all been a waste of time. She told us that unless Tom was prepared to give up drinking and drugs she would be unable to provide any clinical intervention. She referred him to Turning Point and said that she would see him again once he was drug free. Whilst he was willing to try to stop his drink and drug taking, he was unable to engage with Turning Point. Tom had memory difficulties so often failed to turn up for appointments and struggled to comply with treatment due to the nature of his cognitive impairments and mental health issues. He self-medicated because of the intense thoughts and emotions he experienced and without someone helping him to manage those he was never going to be able to stop the drugs. In my view this was a major failing in Tom’s care. We were let down by the NHS and we received no support from social services in fighting his corner. It is my opinion that Turning Point was simply not capable of supporting a man with dual diagnoses and TBI. A specialist treatment service was required that took all of these factors into account (Walker et al, 2007).

By 2012, Tom’s periods of abstinence were becoming few and far between. He was in agony. Tom spent most of his life living in torment psychologically. And in these later years, due to his increasing physical frailty and intense pain in his hip due to osteoarthritis, he was also physically tormented. Tom had always experienced pain as a result of his TBI, something that although highlighted in literature (Nampiaparampil, 2008) was never addressed by
health professionals. Additionally, his increasingly low mood seemed to heighten his
perception of pain (Hoffman et al, 2007). The combination of this chronic pain with the
increasing pain from his hip was unbearable for him. His GP was prescribing pain medication
but between the very real pain he was experiencing and his high tolerance due to his heroin
use, he was eating them like smarties and they were offering very little in the way of pain
relief. It was clear he needed an operation.

Until this point social services had done very little to support Tom. Since 2011, Headway
Somerset had become increasingly concerned about the situation. They informed me that
on numerous occasions they had requested the appointment of a social worker for Tom and
to arrange a case review to look at how he and his girlfriend were living. It took them nearly
a year to even assign someone to his case and even then no direct contact was made.

In early July 2013, Tom was admitted to hospital for his hip replacement surgery but
presented drunk. The operation was cancelled and he was sent away being told he would
need to be clean before he would be allowed back on the waiting list. No time frame was
given to him about how long his abstinence would need to last, or what benchmark he
needed to reach in order to meet the definition of “clean”, and none of this was ever
articulated to me, his advocate. I wasn’t even informed he had been discharged without the
surgery. I spoke at length to Tom’s GP to try and come up with a course of action, but
ultimately we were stuck. He couldn’t stop as he was in so much pain and without the
operation that wasn’t going to change. But without stopping he wasn’t going to get the
operation. And in the middle of all of this it was decided that Tom was a fully functioning
adult who made his own choices so it was his choice all this was happening and the NHS
weren’t prepared to do anything to help. I made it clear to Tom’s GP that the only way he
would get clean was to go into rehabilitation and then straight from there to hospital for the 
operation. But he apparently wasn’t entitled to that support. Social services refused to 
provide support for a referral and eventually Tom was referred back to Turning Point for 
monitoring.

In October 2013, Tom was serviced notice of eviction from his girlfriend’s house for bringing 
drug dealers to her home and allowing them to extort money from her. Social services again 
offered little support. They did not raise any safeguarding alerts regarding the relationship 
between Tom and his girlfriend and they did not provide any support in helping Tom to find 
alternative accommodation.

Just before his eviction, Tom told me he wanted to hang himself but was fearful he didn’t 
have the physical strength to do it, and might leave himself alive but more severely disabled. 
It was at the end of that week that he had a breakdown whilst drunk and presented at the 
hospital asking to be sectioned. That Sunday the crisis team came to his home to do an 
assessment and he was told he wasn’t eligible for treatment because he was simply 
“responding to life events not mentally ill”. Given the vast literature surrounding the link 
between TBI and mental health conditions (Bessell et al, 2008; Bombardier et al, 2010; 
Koponen et al, 2002), the poorer prognosis when associated with substance abuse (Graham 
& Cardon, 2008) and the increased rate of suicidal ideation and suicide risk (Bahraini, et al. 
2013; Felde et al, 2006; Mackelprang et al, 2014), it seemed quite unbelievable to me that 
this was the response.

Tom was eventually evicted from his girlfriend’s home in early December 2013. He had to 
present himself to the housing office as homeless and as there was no intervention from 
social services, he was treated as though he was not a vulnerable adult. During this initial
period, Tom stayed in a few hotels and B&Bs and ultimately with our mother because he kept getting evicted either due to his physical health problems (his seizures meant he was a high risk case) or his drug taking behaviour. He was forced to stay with our mother because no more support was available for him because he had made himself “intentionally homeless” through his actions. There seemed to be little appreciation of Tom’s TBI or his mental health problems. It seemed to me that most of the time the professionals he interacted with simply saw an addict and that coloured their view of him to the extent that they were unwilling to provide him with the support he so desperately needed. Tom eventually ended up at a homeless shelter, adding to the already high rates of homeless people who have experienced TBIs prior to homelessness (Bousman et al, 2010; Hwang et al, 2008; Oddy et al, 2012). At this stage I was still fighting for support for Tom and most importantly some form of Mental Capacity Assessment as it was my firm belief that he was no longer able to make informed decisions for himself. There was also a real issue around safeguarding as he was a vulnerable adult who had now been placed in a dangerous environment surrounded by people who could manipulate and abuse him.

**Tom’s final months**

During the final 6 months of his life, Tom was constantly reporting to us incidents of people stealing his money or phones. There had been a time when we didn’t go more than 48 hours without exchanging texts but during these final months I got used to not hearing from him for days, sometimes weeks at a time. This was usually because his phone had been stolen and he would then have to get a replacement number. It would be ages until I would find out what had happened and make sure he had my number again. During this time his drug and alcohol addictions spiralled even further out of control. After months at the homeless
shelter Tom was found local authority housing in a bedsit in the middle of the worst estate
in the town. He was yet again surrounded by drug dealers and people who would take
advantage of him.

On the 18th June 2014 I saw my brother for the last time. He was in his wheelchair in our
local town. He told me that money had gone missing from his flat and that his card had also
gone. He had given the person responsible his pin number sometime previously to withdraw
some cash for him and now that person had cleared out his account. So he had gone into
town to ask the bank for an overdraft as it was 10 more days until he received another
benefit payment and he had nothing to live on. I don’t know what happened to him in those
last 12 days before he died. I heard nothing from him despite my many attempts.

On the afternoon of the 30th June, my mother, as frantic as I about the situation, emailed
adult social care yet again pleading for help. She impressed upon them yet again how
unstable he was and how she feared he would end up dead in a ditch somewhere. Little did
either of us know that at that point in time that he was in fact already dead. At around 5pm
that afternoon, a policeman arrived to tell us my brother was dead. Tom tied ropes to his
wheelchair and used them to hang himself; a modification to the way he had told me nine
months previously he wanted to end his life. So it shouldn’t have been a shock. But it was.
Not least that he had chosen such a brutal way to end things.

At the end of that week we had to go and pick up what few meagre possessions Tom had
left. Going to his flat was so painful. The place was a tip, with stuff piled up on the bed so
nobody could possibly have been sleeping in it. Yet more stuff was piled up in the shower
cubicle, so it was evident he hadn’t washed for at least as long as he had been in that flat,
which was about two months. And there were used needles everywhere. The worst thing
though was that his wheelchair was still inside the front door, presumably where it had
been left when he died. It was such a surreal feeling seeing this object that I was so used to
being part of him and now viewing it as the device that ended his life.

A few weeks after Tom’s death my mother and I went to a meeting at social services. This
had originally been a safeguarding meeting, but in light of the events it was to review what
had happened. Nobody expected us to turn up. I was horrified during this meeting that
social services turned to us as a family and stated that they had been shocked by the turn of
events as they had no idea this would happen. I think this was the final insult after years of
indifference to Tom’s situation. It was beyond me how anyone could suggest they didn’t see
it coming given the number of phone calls, emails and meetings where we had clearly stated
that this was exactly what would happen.

After Tom’s death, an adult safeguarding review was opened. The report catalogued the
various failings and shortcomings of the care providers and professionals who were
responsible for, and interacted with, Tom during his life. Whilst the report findings are
important, I have taken the opportunity here to outline some of the worst failings from my
perspective as a family member rather than directly reporting the outcomes of the review.
However, following this I have then outlined the recommendations of the adult
safeguarding review along with my own personal recommendations for future practice.

Communication and Information sharing

Since Tom’s TBI, I have been endlessly horrified by the lack of information that is shared
between organisations responsible for his care. Whilst social services often did not have
direct involvement with him, as a vulnerable adult he did fall under their service. Yet
information from organisations such as the NHS regarding Tom’s health, drug use and mental health difficulties rarely seemed to be flagged to them.

Furthermore, we as a family were never informed of any incidents where he presented at hospital. Even after I became his advocate, I was never given this information and was never included in conversations surrounding his care. When I first got to see the findings of the safeguarding review, at points it felt like I was reading about a person I did not know.

Behind the scenes there had been multiple appointments with various people within the NHS and I had not been present and was in fact completely unaware of what had taken place.

Some may wonder why the family of a fully grown man should have such involvement in his day to day care. But the reality of brain injury is that individuals rarely have good insight into their own condition and limitations (Prigitano, 1996). This makes them vulnerable as they will often refuse treatments that they need, neglect their own personal care, and it leaves them open to abuse. As such it is important that family members are kept involved in ongoing conversations about daily care. For example, in 2012, Tom broke his shoulder after falling off his bicycle. I was unaware of this until I found reference to it in the safeguarding review. He should have had his arm in a sling for several weeks whilst it healed, but I didn’t once see him wearing it and neither did my mother. Hence we were unaware of the injury, unable to support his recovery from it, and unable to prevent further injury by helping him to understand why he had been advised to stop riding his bike.

As well as difficulties with insight, Tom also had problems with his memory. This meant that he would regularly miss appointments with his doctor or other health professionals.

Without knowing that these appointments were taking place, there was no way for us to
ensure he attended. Instead, he wouldn’t present at the right time and then be viewed as having dropped out of treatment and be discharged.

Another failing in Tom’s care was a lack of multidisciplinary team working. Tom had a moderate TBI and was a substance user. He was also displaying signs of mental illness. Although, he was assessed by the community mental health team as not having a mental health condition, anyone who knew him will tell you this was a falsehood. During this period each organisation and discrete service wanted to address one issue at a time. Therefore, he was being asked to attend Turning point for his substance abuse, but he couldn’t fully engage in this because he needed psychological support as his addictions were so closely tied up with his mental health problems; he was self-medicating to numb the physical and psychological pain he was experiencing. But he couldn’t get psychological support either because he couldn’t engage in those services until he was “clean”, or because he was viewed as not being mentally ill or suicidal. This was despite having told anyone who would listen that he wanted to end his life. In my opinion, the reason his mental health problems were missed was because his difficulties and behaviour were wrongly attributed to his substance abuse, and that he was never assessed in the presence of an advocate who could speak for him where necessary. His lack of insight into his own condition meant that he would not necessarily have answered honestly during assessment. As highlighted in the literature, individuals like Tom need multidisciplinary team working, a tailored treatment approach and this needs to be guided by an experienced neuropsychologist who can allow and plan for the unique difficulties associated with TBI with regard to lack of insight and poor compliance with treatment (Holloway, 2014; Holloway & Fyson, 2015).
The difficulties in managing the interactions with these various services supporting Tom can be summed up as being a categorical failure to appreciate the biopsychosocial model of health (Engel, 1980). This model highlights how biological and psychological factors can combine with environmental factors to influence individual’s health. In Tom’s case, the neurological damage caused by his TBI, alongside his mental health issues, substance abuse and social deprivation both prior to, and after, his injury led to a distinct vulnerability. At no point was this identified by the organisations working with him. Instead each “difficulty” Tom was experiencing was viewed in isolation, meaning that the organisations involved often missed the bigger picture.

Finally, when providing any kind of long-term care for those with brain injuries, it is imperative that organisations involve the family in trying to understand how much of the exhibited behaviour is related to neurological damage. At no point following Tom’s initial discharge from hospital were we asked to provide any information about what Tom had been like prior to the accident or highlight anything important that might guide his care. Therefore, social services in particular had no benchmark by which to understand his needs and behaviour and often dismissed his unusual appearance and behaviour as being “eccentric” or “anti-establishment”¹ rather than being a symptom of his TBI. Furthermore, as was highlighted in the safeguarding review, Tom had previously experienced three minor head injuries as a child. The literature highlights that even mild TBI can be associated substance abuse, mental health issues and criminality (Sariaslan et al, 2016; Williams et al, 2010), all of which Tom was experiencing pre-injury. This is significant as it suggests he may already have experienced TBI-related complications and then received a significant head injury which may have affected his post-injury prognosis.

¹ These were terms used by professionals about Tom and recorded in the safeguarding review.
The importance of Mental Capacity Assessments

The Mental Capacity Act (2005) provides guidelines to help support individuals with reduced mental capacity to make their own decisions. The Act is designed to protect and empower individuals with various conditions, including brain injury. The premise of the Act states that individuals should be assumed to be capable of making their own decisions unless it is proved otherwise through a Mental Capacity Assessment (MCA). The Act also highlights that just because a decision may be viewed as unwise, that is not necessarily grounds for diminished capacity. Whilst this is an important point, in the case of Tom, he was endlessly making “unwise” decisions both in terms of lifestyle choices and with regards to his care. In the earlier years following his TBI, we were of the opinion that he was still an adult and could still think for himself, so he was able to make most of his own decisions. However, by 2010 he had become so mentally unwell alongside his TBI that it was my firm belief this was no longer the case. Yet at no point was a MCA done. How can one identify whether some has capacity if organisations responsible for their care will not carry out assessments? I believe that because Tom was making unwise day to day decisions and these weren’t necessarily about life or death matters, he was not viewed as requiring a MCA. Yet when his substance abuse prevented him having a major operation to save his ability to walk, this was still not viewed as an issue of capacity.

Furthermore, the Mental Capacity Act states that individuals should have an independent advocate who can support them to make decisions. Tom had an advocate, one that he had chosen himself. Yet I was never privy to any of the conversations regarding his care. It is my belief that had I have been actively included in those conversations he probably would have complied with treatment where he otherwise did not.
The nature of Tom’s dual diagnosis seemed to prevent the appropriate organisations from identifying his need for a MCA. In reality he was a man with a moderate TBI who had a history of undiagnosed mental health problems dating back to pre-injury, and he was a substance abuser. However, it seemed organisations often identified the substance abuse and wrongly attributed his behaviours and physiological condition to that rather than to wider neurological damage and mental health issues.

**Safeguarding and Risk Management**

During Tom’s post-injury years, there were many occasions when safeguarding alerts or risk management assessments should have taken place. Even in the earlier years post-injury where Tom’s behaviour was more stable, Tom should have been treated as a “vulnerable adult”, yet often was not. The Care Act 2014 replaced the term “vulnerable adult” with the definition of “adults at risk”. This change identified that any adult with an illness or condition that may lead to physical or mental impairment should be viewed as “at risk” and that this does not require a formal diagnosis. It is possible that had this change come sooner, Tom would have received greater support from adult social care. However, I admit that I am sceptical. After all, it must be argued that Tom did have a diagnosis – he had a moderate TBI with reasonable cognitive impairment and yet he still was not viewed as “vulnerable” under the old terminology.

Tom’s girlfriend also had a brain injury and was in a wheelchair. For many years he was her main carer. This meant that he was putting constant stress on his already weakened body by performing manual lifting. This should have warranted a risk management assessment but one was not forthcoming. Headway Somerset even alerted social services and requested a
review of their living arrangements once Tom’s physical condition started to deteriorate, but still nothing happened.

There were also numerous times when his substance abuse led him to pass out whilst his girlfriend was in his care. At times she was left on the toilet unattended, or left to sleep in her wheelchair because he couldn’t be woken to help her to bed. Again none of this was flagged as a safeguarding issue for her or him in terms of an indication of his deteriorating mental health.

Subsequently, Tom’s substance abuse reached a point where he was allowing drug users and dealers to frequent the property. His girlfriend was being threatened for money; she had her home alarm system disabled, and her house sprayed with graffiti. There were even claims that Tom himself was extorting money from her for drugs. When adult social care were informed of this, a meeting was held but as Tom and his girlfriend said they didn’t want any help, their social workers left without action. This should certainly have been grounds to initiate a safeguarding alert. A complaint letter I wrote to social services regarding this was left in his file and was never responded to or dealt with.

Tom was later evicted from his girlfriend’s home. During this time adult social care did not provide any support to him with regards rehoming. Tom’s social worker seemed unwilling to provide support and did not respond to our calls for his need for rehabilitation services even after Tom’s own presentation to hospital asking to be admitted for inpatient care. They did not seem to take Tom’s erratic behaviour, or his constant and increasing declarations of suicide, seriously. On one occasion whilst at a Headway Somerset day centre Tom stated his intention to kill himself. Headway spoke to a GP at Tom’s practice who told them it was not their responsibility and to contact his social worker. Social services told them to go back to
the GP as it was not their responsibility either. The GP then told Headway to contact me as Tom’s advocate. This was the only time my role as an advocate had been acknowledged and in this instance it was wholly inappropriate. One or both of those organisations should have raised a safeguarding alert, but neither did.

During the last months of Tom’s life, he was living in a homeless shelter and then a local authority bedsit. In both cases he was living in close proximity to individuals who were taking advantage of him. Yet, it still took five months before any kind of safeguarding alert was made. Sadly this was far too late. Meetings took place between all relevant agencies and Tom underwent a risk assessment about a week before his death. Again I was not present at this assessment and was unaware it had happened until after his death. The paperwork from it shows that Tom did not fully highlight all the difficulties he had been experiencing, or the personal and financial abuse he had been experiencing. He also refused rehabilitation services. In my opinion this was partly because there was nobody there to support him through that decision making process, and because by that point he had given up hope. He needed someone in that meeting to explain things to him and be there for him. Perhaps then things would have been different.

**Recommendations for future Practice**

To conclude, I have outlined below a brief summary of the main recommendations made by the safeguarding review conducted in to Tom’s care and subsequent death. Below these recommendations, I have provided my own insight and comments on why they are fundamentally important for shaping current practice:
• Tom’s case study features in sector-led and multi-agency training…and that multi-agency work with individuals with complex needs is shaped by shared goals and clear leadership.

• A learning event should be hosted concerning Tom and others currently known to Headway Somerset.

• The fact of a person’s TBI and mental capacity is foregrounded in all professional assessments and referrals and that family involvement is prioritised with a view to understanding the continuities and discontinuities and the unpredictable and complex process of reconstructing the self which arise from such critical injury.

• Public Health, Somerset County Council, and NHS commissioners should set out how local practice and priorities match good practice concerning the support of people with brain injury, dual diagnoses and the expectations of the National Suicide Prevention Strategy for England.

• Homefinder Somerset and Housing partners identify how tenants with extensive support needs, including those with acquired brain injuries, may access supported housing.

• The review is shared with Headway UK for dissemination beyond Somerset to stimulate debate.

These recommendations provide concrete actions following the safeguarding review. Below I have tried to elaborate on some of these by thinking about the issues that would have directly improved care for Tom.
Echoing the recommendations above, it is my belief that the NHS and social services need to work together to ensure information about vulnerable adults is routinely shared for the better management of their care. They often have complex needs and must be looked after by specialist multidisciplinary teams (Headway, 2014). Organisations need to take a more active role in case management for individuals with TBI. There needs to be one individual accountable for information sharing and to act as a conduit for support and care for individuals and their families and to provide better access, and signposting, to services. Adult social care need to undertake regular specialist assessments to monitor the health and wellbeing of individuals with TBI (Holloway & Fyson, 2015). It is also imperative that Health and Social work professionals receive better training in understanding the difficulties associated with TBI, particularly with regards to mental capacity and dual diagnosis. Specifically, more awareness is needed of the co-occurrence of TBI alongside mental health problems, suicidality and substance use.

Routine involvement of families is crucial in the care of those with TBI to circumvent the difficulties caused by lack of insight. Lack of insight is associated with diminished capacity. Furthermore, because individuals following TBI have residual intellectual capacity, there is a very real danger that they can present as being capable of making complex decisions (Holloway & Fyson, 2015). The advocacy role must be taken more seriously and routine contact made by all organisations. Family and advocate involvement is crucial to ensure cases of diminished capacity are not overlooked.

Conclusions

Although none of what is highlighted above will bring back my brother Tom, it is my hope that this case study can provide information that will help to save the lives of other
individuals with TBI in the future. By providing more consistent sensitive care with a better understanding of the specific difficulties TBI presents, others in the future can receive the kind of support that my brother and our family needed. To conclude, I would like to remind professionals that brain injury is something that can happen to anyone. It is possible that this very day you yourself may experience such an injury, or someone you love. With that in mind, it is the responsibility of all to ensure that individuals with brain injuries receive the kind of care and support that we ourselves would like to receive.
References


