2 1 Monday, 24th February 2020 1 189 witnesses. I have personally read over 2 (10.32 am) 2 a thousand statements, and that's an ongoing 3 3 SIR BRIAN LANGSTAFF: Ladies and gentlemen, let me process. I undertook to read every single 4 tell you what this week has in store for us. 4 statement no matter how many there were, and 5 5 First, let me put it in context. At the start are, or will be, and I am doing. But it was 6 of this Inquiry, I promised that I would hear 6 obvious from the start that there were many who 7 from those infected and affected both first and 7 wanted their voices to be heard but did not wish 8 last. I've almost, but not yet quite, honoured 8 or did not dare either to give evidence orally, 9 9 that promise so far as putting you first is or to make a written statement knowing that it 10 concerned. 10 would be disclosed, even though their identity 11 11 could be anonymised. This week is something of a stepping stone, 12 a transition between listening to the testimony 12 So this Inquiry did what I believe no other 13 of those who have suffered, and then from June 13 Inquiry has yet done: it found a way of giving 14 onwards this year, taking the evidence of those 14 a voice and providing a public platform for this 15 who are in positions of responsibility, as 15 group of people in a way that can be taken into clinicians, administrators, politicians, 16 16 account. It asked trained experienced 17 suppliers, and others, when choices were made 17 interviewers to speak to anyone who wished it, 18 which are said to have led to that suffering, or 18 and to present a report which drew on all the 19 which failed to deal with that suffering, 19 accounts given to them, so that it presented 20 properly. 20 a comprehensive overall picture of what they had 21 Within this week of transition, today marks 21 been told. The Inquiry calls them 22 a stepping stone of its own. I said I'd not 22 intermediaries because that's what they are. 23 quite honoured my promise so far as hearing 23 They provide another way, in addition to written 24 24 first from people who have been infected or statements and oral testimony, by which the 25 25 affected, I have heard oral evidence from voices of people and their families can be 3 4 1 heard. 85 people have taken advantage of this 1 what has been said. The link between the 2 2 evidence of the intermediaries and the panel of so far After the intermediaries have presented 3 3 psychosocial experts is a close one. For 4 their findings, then my promise will have been 4 instance, one expert conducted research projects 5 fulfilled as far as those from whom I said 5 focusing upon recording the life histories of 6 I would hear first is concerned. The evidence 6 those with both haemophilia and HIV, and their 7 7 of the intermediaries leads on neatly, later families. In doing so, she helped capture the 8 today, to the evidence of experts -- there are 8 experience of roughly the same number of people 9 9 seven in all, counting both today and again as did the intermediaries. You can see 10 tomorrow -- who have spent their professional 10 how the evidence follows on naturally. 11 lives at the highest levels of psychology and 11 But the evidence may achieve more than 12 sociology, studying how people react to disease, simply telling you people's stories. Many of 12 13 particularly disease which is life shortening or 13 those who have suffered tell me they want to help to avoid anything like what happened to 14 life threatening, and affects every aspect of 14 15 a person's life, their relationships with their 15 them happening to others in the future. And 16 partners, family and friends; their schooling; 16 these experts will explain how the accounts of 17 the work they can do; their housing; their 17 those from whom we have heard fit with the 18 finances; their ability or desire to bring 18 learning derived from the experiences of others, 19 children into the world; their need for care; 19 reflected in expert journals and in their own 20 their trust of those to whom they may still have 20 extensive studies, and what is known about the 21 to turn if they seek treatment. 21 best ways to reduce the risk of any repeat in 22 22 the future. That's something I want to consider Although they have different roles, the 23 23 intermediaries report what is being said to them closely. 24 without analysis or comment and the psychosocial 24 The evidence of the psychosocial experts 25 experts provide analysis of and commentary on 25 will finish tomorrow. It forms a natural bridge

5 6 1 between the lived experience of those who 1 and hepatitis can be reduced. 2 provided evidence in any of the three ways 2 Let me be very clear about what the clinical 3 3 I have mentioned and the clinical experience, experts have not been asked to deal with. They 4 research, knowledge and expertise to be given by 4 will not say what they think was right or wrong 5 5 about what happened historically. They will not the clinical experts from whom we shall hear on 6 Wednesday about hepatitis, Thursday about HIV, 6 say what happened and when back then, except 7 and Friday about bleeding disorders. 7 only by way of background to explaining today's 8 I promised that this Inquiry will be open 8 medicine. What happened and when are matters of 9 and transparent. And what the Inquiry has asked 9 fact, upon which the Inquiry will draw its own 10 10 the experts has been disclosed. How they have conclusions of fact. So please, don't expect 11 answered it has been disclosed. And further 11 any questions to be asked about that at this 12 supplementary questions prompted by core 12 stage of the Inquiry. 13 participants through their representatives are 13 Now, counsel to the Inquiry will explain the 14 all open to view. 14 mechanics of what is proposed for today and for 15 Why do we need to hear this expert evidence 15 the rest of the week. Before we start this 16 now? Well, the answer is threefold. First, to 16 morning's evidence from our three 17 provide a solidly based medical understanding 17 intermediaries, Jackie Wilson, Kay Durrant and 18 that will inform both the questioning of other 18 Pam Allen, who wish to be known as Jackie, Kay 19 witnesses later this year and the report of the 19 and Pam. 20 Inquiry. 20 Ms Richards. 21 Second, to let you know what the latest 21 MS RICHARDS: Sir, as you've indicated, we will be 22 state of knowledge is about conditions; many of 22 starting this morning with the intermediaries 23 you will know only too well by experience. 23 and Ms Scott will be taking them through their 24 24 Third, so the wider public can hear, and as evidence. 25 25 they hear, that ignorance about HIV infection This afternoon we will start to hear from 7 8 1 the evidence of the psychosocial group. There 1 Dr Ian Williams, Dr David Johnston, 2 2 will be five members of the group giving Professor Graham Cooke and Ms Sian Edwards. 3 evidence this afternoon, Professor John Weinman, 3 Dr Gareth Tudor-Williams is probably going to be 4 Professor Myfanwy Morgan, Dr Nicky Thomas, 4 unable to join us on Thursday. 5 Dame Lesley Fallowfield and Ms Sian Edwards. 5 Then finally, on Friday, we will hear about 6 Professor Deborah Christie is unable to attend 6 bleeding and blood disorders from 7 7 Dr Oliver Tunstall, Dr Richard Gooding, today but will be attending tomorrow morning, 8 and Dame Theresa Marteau will also be unable to 8 Dr Sara Marshall, Dr Mallika Sekhar, 9 9 attend today but will be available tomorrow Professor John David Edgar and 10 afternoon. 10 Professor Jürgen Rockstroh. Professor Rockstroh 11 Sir, as you've indicated, one member of the 11 will be joining by video link in the morning 12 psychosocial group, Ms Sian Edwards, was closely 12 only. 13 involved with the HIV and haemophilia life 13 Sir, core participants have already, through history project, and we will be taking advantage 14 14 their recognised legal representatives or 15 of her presence here to ask her to tell us about 15 otherwise, put forward suggested questions for 16 that, in the course of either this afternoon or 16 the expert groups arising out of their reports. 17 17 Ms Scott and I will not be necessarily be asking tomorrow morning. 18 The psychosocial evidence will continue into 18 every one of those questions, some of them later 19 tomorrow and then Wednesday's focus will be 19 in the inquiry, some are perhaps more 20 hepatitis and we will be hearing from 20 appropriately directed to other expert groups, 21 Professor Graham Cooke, Professor John Dillon, 21 such as the ethicists, and other questions that 22 Dr Kate Jeffery, Dr Scott Jamieson and Dr Aileen 22 were suggested ask experts to address matters of 23 23 Marshall. fact which are, for the reasons you've 24 On Thursday the evidence will focus on HIV, 24 explained, sir, ultimately for your decision. 25 and we will hear from Professor Jane Anderson, 25 But we will break towards the end of the

9 1 evidence of each expert group, so that core SIR BRIAN LANGSTAFF: Thank you very much. May we 2 participants can have the opportunity to raise 2 then have Jackie, Pam and Kay, please. 3 with Ms Scott and myself further questions that 3 KAY DURRANT, sworn 4 4 JACKIE WILSON, affirmed they would like to be asked. 5 5 PAM ALLEN, affirmed Sir, finally in relation to the mechanics of 6 this week, the Inquiry has asked those 6 Examined by MS SCOTT 7 responsible for the NHS in the four parts of the 7 MS SCOTT: Can we start by each of you giving an 8 United Kingdom, Northern Ireland, Wales, 8 introduction to your professional background and 9 9 Scotland and England, for information about the explaining the skills that you brought to the 10 availability of psychological support and the 10 intermediary role. 11 arrangements made for the commissioning and 11 Pam, do you want to start us? 12 funding of psychological support, and for PAM: Yes, I'm happy to start, thank you. 12 13 information about current Hepatitis C care and 13 A qualified social worker since 1979, which 14 treatment and in particular, issues about scans, 14 clearly put me in a position of dealing with 15 blood tests, checks and monitoring. 15 people who've experienced trauma via their own 16 The Inquiry has now received witness 16 abusive situations or their experiences, 17 statements that address those issues. They will 17 together with lots of experience of people with 18 be disclosed or should be disclosed, I hope, to 18 bereavement and loss in various aspects of their 19 19 life. I also worked for the court during that core participants in the course of today, and in 20 the course of this week, I will explain and 20 time preparing reports in a formal setting. 21 summarise what those statements say at an 21 MS SCOTT: Kay, do you want to go next? 22 appropriate stage on one of the days. 22 KAY: I'm a former senior detective, having served 23 Sir, that's all I needed to explain by way 23 for 25 years in the police service, and done 24 of mechanics, and we're ready now for the 24 lots of work around trauma in the public 25 evidence of the intermediaries. 25 protection arena, in particular around family 12 11 1 liaison officer roles in the homicide MS SCOTT: The chair in his introduction said that 1 2 2 investigations. I then latterly, in 2012, you'd spoken to around about 85 people; is that 3 commenced training as a psychotherapist, and I'm 3 right? 4 involved in private practice, and again, that 4 KAY: Yes, yes. 5 has enhanced my knowledge with regards to 5 MS SCOTT: Jackie, can I ask you, when did you carry 6 dealing with those with trauma. 6 out these interviews and over what time period? 7 MS SCOTT: And Jackie, do you want to? 7 JACKIE: Over around a 12-week period last year, 8 JACKIE: Yes, I'm also a social worker, I've been 8 largely between the end of January and the 9 9 qualified for 38 years and, like Pam, I've beginning of April 2019. 10 worked across a number of sectors in the family 10 MS SCOTT: What was the geographical spread of those 11 courts in local authorities, but also in 11 that you spoke to? 12 hospitals as well, in paediatrics. And I'm also JACKIE: We saw people in England, Wales and 12 13 just coming to the end of 4 years of training as 13 Scotland, and Northern Ireland. MS SCOTT: In page 3 of your report, you say this 14 a therapeutic counsellor as well 14 about the range of experience of those that you 15 MS SCOTT: Before you talk about the substance of 15 16 the report, and what you were told by those you 16 interviewed. Six had been infected with HIV, 17 spoke to, can we just establish your method of 17 50 had been infected with hepatitis C, 11 had 18 working? 18 haemophilia, three had thalassaemia, nine were 19 Can I start with you, Kay. Approximately 19 infected through routine surgery, 12 were 20 how many visits were carried out between the 20 infected through emergency surgery, eight were three of you? 21 infected in connection with childbirth, and one 21 22 We visited between 50 and 60 homes or places 22 person had been infected by their partner. KAY: 23 23 to meet with individuals and family members, and Does that summarise that range of 24 the age range was meeting people between the 24 experience? 25 ages of 30 and up to the age of around 80. 25 KAY: (Witness nodded)

	13	14
1	MS SCOTT: Jackie, can I ask you then about the	1 and some of our longer ones were over 5 hours.
2	interviews themselves. Where was it that you	2 So we think probably an average of three, three
3	saw people?	3 and a half hours.
4	JACKIE: We saw people mainly in their own homes.	4 MS SCOTT: Pam, can I ask you then about the report
5	We gave people the choice of where they would	5 writing. In terms of communication that you had
6	like to be seen. Most people chose to be seen	6 with those you interviewed after the visit, what
7	in their own homes. Some people, for reasons of	7 kind of communication did you have with them?
8	confidentiality, didn't want to be seen at home,	8 PAM: I suppose I would start by saying that all
9	because of other family members being aware of	9 our interviews were very emotionally charged and
10	our visit, and so sometimes we saw people in	10 were could be quite traumatic in some
11	conference centres, booked rooms, etc. Some	11 circumstances, because this was the often the
12	people actually wanted to be seen quite some way	12 first time that the people we were interviewing
13	away from their own home town as well. So we	13 had told their story.
14	saw a small number of people in those	14 We established that the groundwork there,
15	circumstances as well.	that we would produce a report, which would be
16	MS SCOTT: And did you do the visits alone, did you	a joint report, which was confidential, and we
17	divide them up between you?	17 would also agree on which elements of that
18	JACKIE: We did we largely went alone, except	18 report people were then happy that were included
19	that Kay and Pam did the Northern Ireland visits	in this report that we produced, for the
20	together.	20 Inquiry.
21	MS SCOTT: How long would a visit take on average?	21 So the methods that we used, we invited
22	JACKIE: There was a range of visits. We weren't	people to tell us their story. It was their
23	time limited at all. Our remit was we're here	23 time to tell us their story, and walk through
24	to listen to whatever people wanted to say to	their story. We took detailed notes at that
25	us. None of the visits were less than 2 hours,	point. We then went away, developed the report,
1	15 which we sent to our person. They added,	16 1 their identity private?
2	subtracted, changed, got to a point where we	2 <b>PAM</b> : Absolutely, and that of their families,
3	both comfortable. So they were jointly authored	3 obviously, which is still around stigma and the
4		5 Obviously, which is still around stignia and the
h	reports, to the point that we put both our names	4 fear of that stigma. And also confidence.
	at the bottom of the reports. So it's an agreed	<ul> <li>fear of that stigma. And also confidence.</li> <li>I think they felt that having an interview in</li> </ul>
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	17		18
1	experiences of all of the 80 people that you	1	most traumatic times of their life, and some of
2	spoke to, but am I right in understanding that	2	that, many of the families, it's still continues
3	you have taken into account everything that	3	with infections today, and those that have lost
4	you've been told in your report	4	loved ones, and what was definitely needed was
5	KAY: Yes.	5	that ethic of care in terms of how we obtained
6	MS SCOTT: writing and your preparation for	6	that information. And I think we all agree it's
7	giving evidence today?	7	one of the most significant things in our
8	PAM: Yes.	8	professional careers that we've contributed to,
9	JACKIE: Yes.	9	and we're honoured to do that and want to do the
10	MS SCOTT: Can we start, then, by you set out	10	best that we can today to represent all our
11	both at the beginning and the end of your report	11	families.
12	that you were all humbled by the experience of	12	MS SCOTT: Is there anything anyone wants to add to
13	speaking to these 85 or so people. Is there	13	that?
14	anything you would like to say about the impact	14	JACKIE: I think we were really struck by people's
15	on you of having undertaken this exercise?	15	dignity. The dignity that people showed, and
16	KAY: I mean just to expand on that a little bit,	16	their strength in managing these situations over
17	I think all of us felt completely humbled and	17	years and decades in some instances. And we
18	privileged that families were telling us some of	18	felt that as Kay said, it was a privilege
19	the most painful experiences in their lives.	19	we felt that we saw the best of humanity.
20	And whilst we are professionals, you can't not,	20	PAM: I think what I'd add to that is what we've
21	on a personal level, feel sad yourself, but	21	stated in our report, that the people we met all
22	obviously our professional hats were really	22	said, "We're the lucky ones, we're the lucky
23	important but it was also really important to	23	ones, because we're still here." And that, to
24	show that kind of level of impact of hearing	24	hear that from people who have experienced such
25	that, and families were telling us some of the	25	trauma and such massive difficulties in their
	19		20
1	lives, automatically humbled us. Most of the	1	consultant. And this was in close proximity,
2			
_	interviews had tears, and that was the tears of	2	the hospital and the college he was at. And he
3	interviews had tears, and that was the tears of those people who were interviewing, and our own.	2 3	the hospital and the college he was at. And he had a phone call that morning to say can he go
3			
3 4	those people who were interviewing, and our own.	3	had a phone call that morning to say can he go
3 4 5	those people who were interviewing, and our own. It was a very consolidating experience having	3 4	had a phone call that morning to say can he go down to see his consultant, go down to the
3 4 5 6	those people who were interviewing, and our own.  It was a very consolidating experience having interviewing these people. We made an	3 4 5	had a phone call that morning to say can he go down to see his consultant, go down to the hospital, which he did, without talking to his
3 4 5 6 7	those people who were interviewing, and our own.  It was a very consolidating experience having interviewing these people. We made an attachment within those hours that we spent with	3 4 5 6	had a phone call that morning to say can he go down to see his consultant, go down to the hospital, which he did, without talking to his parents or anyone else in his family.
3 4 5 6 7 8	those people who were interviewing, and our own.  It was a very consolidating experience having interviewing these people. We made an attachment within those hours that we spent with them, which is probably quite different to other	3 4 5 6 7	had a phone call that morning to say can he go down to see his consultant, go down to the hospital, which he did, without talking to his parents or anyone else in his family.  He went down to see his consultant to be
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3 4 5 6 7 8 9	those people who were interviewing, and our own.  It was a very consolidating experience having interviewing these people. We made an attachment within those hours that we spent with them, which is probably quite different to other things that we've done in our careers.  JACKIE: Mm, mm.	3 4 5 6 7 8 9	had a phone call that morning to say can he go down to see his consultant, go down to the hospital, which he did, without talking to his parents or anyone else in his family.  He went down to see his consultant to be told that he was HIV positive and that the incubation period was probably 5 years, he'd had
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22 1 status. Certainly a story I will never forget. 1 old son has HIV". 2 The next young man was 19. He, similarly 2 Desperate for more information, the parents 3 3 haemophiliac, was told he was HIV positive and contacted the hospital; this young man was 4 had been for 3 years. Again, the incubation 4 haemophiliac and they had specialist 5 period, and he do thought that he'd got 5 professionals that they dealt with. And the 6 approximately 2 years to live. 6 consultant was on holiday for another two weeks, 7 He described being in a state of denial, and 7 and this time for them was just horrendous in 8 just taking himself off. He couldn't actually 8 terms of getting that in the manner that they'd 9 remember where he'd lived for a few years after 9 had that information delivered, and then having 10 that. He'd wandered and not been part of his 10 to wait for a further two weeks to have any 11 family, got odd jobs, and it took him a good few 11 conversation with a consultant. 12 years to actually come to terms with what had 12 And this family later found out that the son been said and to seek out treatment. He, again, 13 13 had been infected some years earlier, this was 14 is in his fifties now. 14 backed up by medical notes, but in particular, 15 MS SCOTT: Kay, you spoke to a family who were given 15 there was a big thing for them because they were 16 diagnosis by letter. Can you tell us what you 16 hearing at the time that if you could survive 17 were told by them? 17 HIV for more than 2 years, then you were likely 18 KAY: Yeah, both this case and another case that 18 to live longer, and for desperate parents, they 19 19 were really trying to get information and that I'm going to talk about were two families where 20 I met the parents of children who had HIV. The 20 was very difficult for them. 21 first one was a young boy who was 11 years old, 21 The second family was a couple's again young 22 22 and the devastating news was not delivered in son who was wrongly diagnosed with haemophilia 23 person but a letter that they received one day 23 and later in years he was found to not be 24 24 and just went to the front door and opened this haemophiliac. He was 3 years old, and later in letter and in this letter it said, "Your 11-year 25 25 his life, when he was about seven, they found 23 24 1 out that he was HIV positive. And again, it 1 infection and diagnosis with hepatitis C. 2 took his parents two months to get the results, 2 JACKIE: We saw a large group of people in 3 and again, that pain and agony of waiting and 3 respect of this issue and we've been able to 4 then obviously receiving the news at the end 4 find three key themes that we want to speak 5 that their son had HIV. The commonality again 5 about today. 6 between the two families is when their children 6 The first one was when people presented with 7 7 were informed that they were HIV, which for both symptoms, there was a reluctance and 8 families were around the age of 13 or 14, where 8 a significant delay in professionals 9 9 they'd started to form relationships and both of acknowledging that these might be as a result of 10 the young boys were getting girlfriends and the 10 hepatitis C, so a lack of investigation, and an 11 parents had to then share the information with 11 explaining away, really, of symptoms, 12 their sons, and, you know, responses such as "Am particularly when people presented with the very 12 13 I going to die", and having to deal with that 13 significant fatigue that goes with it so people were describing -- they said it was because of 14 with a teenager, were extremely traumatic for 14 15 both families. 15 their stage in life, so if they were young 16 16 children, women who had just had had babies, if And to this day, both of those boys are 17 still alive, living with HIV, and one of the 17 they had demanding work roles, et cetera, there 18 parents says every time a telephone rings, 18 was a almost a default mechanism of kind of 19 "That's my life", in terms of "Am I going to 19 explaining it away, and sometimes that went on 20 find out that he's died?" And it's just that 20 for years into more than a decade, and people 21 constant fear of their child dying, even though 21 spoke about all that time, the disease was 22 we're a number of years ahead, is still as 22 silently progressing within them. Obviously 23 powerful today as it was then. 23 they had concerns about infecting others and 24 MS SCOTT: Jackie, can I ask you then to start us 24 we'll come on to speak about that later. 25 off on the next theme in your report, which is 25 The second theme was about some real

25 1 insensitivity in the way that people were 1 Very few people were given detailed 2 informed about their hepatitis C status. 2 information about hepatitis C, we found, when 3 3 they were first diagnosed. And many people Then the third area was once the hepatitis C 4 was recognised and confirmed, a real reluctance 4 described having to do that research for 5 to acknowledge that it might have come from 5 themselves, particularly if they had any 6 infected blood or blood products. So, for 6 specialist issues. And some of their GPs 7 example, one chap who was -- had a transfusion 7 actually acknowledged that and saw them as 8 as a result of an injury, late in his life, in 8 experts by experience, and were grateful for the 9 9 his sixties, and then developed the hepatitis C. help that they were given. But it was 10 He'd been in the army during the war and the 10 unsettling for people to think that their 11 doctor asked him if he'd been a drug user or 11 clinicians didn't have the level of experience 12 shared to needles in the army in the 1940s and 12 that they wanted them to have. 13 he was horrified by this. It felt a real slur 13 Lots of people described themselves being 14 14 asked about their lifestyle, particularly in 15 Lots of people described being told they 15 relation to alcohol use. Many were asked 16 must have caught it abroad. One woman told me 16 whether they'd used drugs and whether they'd had 17 that her late husband was told by his specialist 17 contact with sex workers, which was terribly 18 that he was imagining the symptoms, and when she 18 demeaning for them. 19 accessed his records after he'd died, the record 19 In terms of how people were told, one man 20 said, "This is a miserable man, always 20 told me he had haemophilia, and he went to his 21 complaining", and this man had cirrhosis by the 21 regular haemophilia clinic with his wife and two 22 22 time he was diagnosed. small children and was told in the room without 23 23 any warning that he had hepatitis C and that he One woman was told by a doctor who didn't 24 24 examine her physically, and didn't do any tests, had 3 years to live, and he had his children on 25 25 that she had ME rather than hepatitis C. his knee, a three-year-old, and his wife had a 1 27 28 1 year old baby. 1 brings to the fore the trauma of losing her 2 2 Another man who was 23 at the time had baby, losing her child, and that she didn't give 3 thalassaemia, he was told by his consultant that 3 consent to any of the treatment at that time. 4 he had tested positive for hepatitis C and this 4 Her husband clearly felt that he needed to 5 conversation took place in a corridor in the 5 make that decision because her life and the life 6 hospital. He'd not been asked whether he'd 6 of the baby were both hanging at that point and 7 7 consented to the testing being carried out and he needed to make a decision, so the trauma of 8 there was no discussion about treatment, as with 8 having to talk about how she became infected and 9 9 Pam's description, the only advice he was given the loss of a child is clearly something which 10 was that he shouldn't have unprotected sex. 10 is unimaginable. 11 And again, this was discussed in the 11 Moving on from that, this lady then 12 corridor. There was no other description at experienced some of what Jackie has just 12 13 that time of the implications of hepatitis C. 13 referred to for years, the fogginess, the depression, the weight loss, the fatigue, and 14 Pam's got some more examples. 14 15 PAM: Yes, I met a lady who was infected at the 15 didn't know until another 20 years on that she 16 point that she had needed to have an emergency 16 actually had hepatitis C, and how that had been 17 caesarean. Child was at 25 weeks, and placenta 17 contracted. 18 was ruptured. Her husband had made the decision 18 She also -- and we go on to talk about it 19 about needing the caesarean and following that, 19 later on in the report -- was a blood donor, 20 obviously she needed blood. She was 20 because she was grateful that she'd actually 21 haemorrhaging, and this was the point that she 21 been given blood, which had kept her alive. She 22 became infected. She didn't know that for 22 went on to be a blood donor, and we'll talk 23 23 20 years, and 20 years on. about that later on and the impact there of the 24 The point of this here is twofold, that 24 guilt of that instance as well. 25 talking about how she became infected clearly 25 And the late diagnosis that I met, like

30 1 Jackie and Kay, a number of women who all 1 that she'd been diagnosed 14 years previously, 2 experienced these symptoms of fatigue, poor 2 and it was actually in the notes, and the GP had 3 3 memory, severe aches and pains, confusion and taken the decision not to inform her or the 4 saw GPs over many years who talked about their 4 family of the diagnosis based on the patient's 5 5 busy lives and that there were women who were mental health. 6 working, sometimes had children, and weren't 6 This lady did have depression, which is no 7 dealing with the stress of their lives very 7 shame at all, and she had capacity and she had 8 well, and it was often into 20, 30 years before 8 that choice taken from her for 13 years; and she 9 they were diagnosed. 9 felt very, very strongly that this wasn't 10 a decision for a professional to take away from So during that period of time their 10 11 confidence, their lack of self-esteem, their 11 her. That should have been hers. And she said 12 feeling of vulnerability, had been layered upon 12 to me, "I should have been told, because I could 13 layered during that period pre-diagnosis. 13 have infected by family." 14 I think Kay has got more examples there, 14 The impact that had on her psychologically, 15 15 finding that out, 13 years later, I can't too. 16 KAY: I just wanted to continue the theme around 16 describe, you know, how she reflected that day. 17 the late diagnosis of hepatitis C. One of the 17 She was a lady then in her eighties and it had 18 families I met a lady who was infected with 18 had such a profound impact on her life, knowing 19 hepatitis C and her two daughters. Sadly the 19 that these 13 years she could have been 20 lady passed away last year so isn't here to 20 responsible for hurting somebody else. 21 obviously hear this evidence being given today. 21 Her actual daughter said to me, "You know, 22 22 I think it's really important that I reflect it's disgusting and it makes me angry that they 23 something she said that day as well to me. But 23 didn't tell her. It was irresponsible and it 24 put my mum, the family and the health 24 one of the issues for her was that her family 25 25 had requested the medical records and found out professionals at risk". 31 32 1 Just to touch on two other cases as well, 1 progress of the illness was described to her, 2 family cases that I heard of, was where families 2 she was given a flow diagram which had been 3 had pre-existing conditions, and yet didn't find 3 produced by the British Liver Trust and she kept 4 out until 2004 and 2009 and yet they'd been 4 it and I saw that. It looked to be part of 5 being having all these blood tests being taken 5 a Powerpoint presentation that has been prepared 6 for years and years and years and one of the 6 for professionals, and the diagram showed the 7 7 ladies found out, because she was progressing progression of hepatitis C and its implications 8 through to IVF procedures and found out in 8 and it showed a life expectancy of 30 years. 9 a private process that she had hepatitis C, not g There was an arrow that went across with 10 through any other procedures, and likewise, for 10 "Diagnosis" with 30 years and "Death" being at 11 the other gentleman as well. He had 11 the end of it and it showed the progression, 12 a pre-existing condition that meant that he from potential infection to cirrhosis and liver 12 13 regularly had to have blood tests but again, 13 cancer to the end stage. 14 didn't find out until 2004, and they have real 14 At this point it was 26 years since the 15 questions around, you know, even if it was in 15 woman had been infected, and she said that she 16 the early nineties, where hepatitis C was, you 16 was left to make the calculation herself. She 17 know, there was a test available, why did it 17 said, "I had to do the maths", which was that 18 take so long for us to find out, and the 18 she had got less than 4 years to live, and she 19 implications of us having that for so many years 19 did unfortunately pass away within that period. 20 was so important for us to know, and both very 20 MS SCOTT: Pam, I think you also spoke to a number 21 of people who were angry that there hadn't been 21 angry about that. 22 22 a re-call programme that could have identified JACKIE: I met with the family of one infected them earlier. 23 woman, who, when she was eventually diagnosed 23 24 after her diagnosis, she went to an appointment 24 PAM: Yes. 25 with consultant hepatologist, and the way the 25 MS SCOTT: Do you want to speak about that in a bit

33 1 more detail? 1 same symptoms didn't know about each other, and 2 PAM: Yes. 2 they saw that HIV and hepatitis C was something 3 3 JACKIE: Yes, we will speak about that later. Do that was generally about people who'd led 4 you want to talk about that now? 4 a different type of life, or who indeed had 5 PAM: Yes, I will give a couple of examples here. 5 illnesses such as haemophilia. They did not 6 The first person I visited was a woman who had 6 understand that there were women out there with 7 had blood transfusions in 1973. But only been 7 hepatitis C in similar circumstances. That in 8 diagnosed with hepatitis C 3 years previously, 8 itself, I started to say to these women, "There 9 9 and she started to tell me her symptoms and what are other women with your experiences", and that 10 10 life had been like during those intervening brought some comfort. 11 years by way of her health. It was the first 11 The second lady I saw, who'd had blood 12 time I'd heard this, and took my notes, and she 12 transfusions in 1985, the way she found out 13 was a very, very brave lady, because she was 13 about her hepatitis was again, very traumatic. 14 very unsure about contributing to the Inquiry, 14 She'd spent, as I say, 30 years, this lady, 15 I'd had telephone conversations with her before 15 experiencing these dreadful symptoms that were 16 I met her, and indeed, once she started talking, 16 very debilitating for her. Then she had what 17 it was like the floodgates had opened. 17 she saw as routine blood tests as she was 18 She, as I say, gave me a perfect account of 18 entering menopause, possibly, went to the 19 the physical symptoms and emotional symptoms and 19 hospital for her results, and the consultant's 20 how her life had changed. 20 line of enquiry entirely confused her, asked if 21 The next lady I saw, I knew what she was 21 she was or had been a sex worker, asked if she'd 22 22 going to say when she started to talk to me ever taken any intravenous drugs or snorted, and 23 about her symptoms. I knew what was coming and 23 she said she didn't even understand what 24 "snorted" meant. 24 throughout my interviews, I certainly understood 25 25 Then asked her if she'd been involved in that these women who had all experienced these 35 36 1 alcohol, whether her partner had been involved 1 clearly something that she could never recovery 2 2 with sex workers. She asked if he could look at from his notes to check that he was seeing the right 3 3 JACKIE: The lady that I spoke about earlier who had 4 person, had he got the right name, she was so, 4 seen the Powerpoint slide, she was a woman who 5 so confused about the line of questioning, and 5 kept -- left a written testimony of her 6 highly distressed at the line of questioning. 6 experience of the infection, and in this she 7 7 At some point, then, she was trying to think wrote, and I want to quote this, she said: 8 about what possibly could have happened. It was 8 "I feel that my country knew where they were 9 9 actually the same hospital that she was getting sourcing blood from, and when they had the 10 these results, where she'd had a blood 10 opportunity to correct this terrible disaster by 11 transfusion some 30 years since with an ectopic 11 completing a re-call programme, they failed to 12 pregnancy and it was only at that point that the carry it through. This was proposed in 1985. 12 13 consultant was able to change their approach to 13 If I had been traced and tested in 1985, then 14 her. She'd gone on this visit for her results 14 the virus would have been more treatable, maybe 15 alone because she expect the results of 15 even clearable, and today I would not be left 16 a standard blood test, so was entirely and 16 with the cirrhosis of the liver which could lead 17 17 to liver cancer. I'm left with a possible death deeply traumatised. 18 Throughout this period when she had felt 18 sentence. It's destroyed all of my plans for 19 very vulnerable and weak because her health had 19 the future, and I've lost my ability to dream." 20 been so poor at least having an explanation 20 This lady did unfortunately pass away 21 about why her health had been so poor and her 21 relatively soon after that. 22 22 MS SCOTT: That anger at there being no recall emotional wellbeing had been so poor, had at 23 least explained to her that it wasn't about her 23 program, is that something you heard from other 24 difficulties but actually it was a physical 24 people that you see. 25 illness, but the way that this was delivered was 25 KAY: It is.

37 1 JACKIE: It was a really strong issue. 1 appointment to go and donate blood, which he did 2 MS SCOTT: Before we move on to the next theme, is 2 every year, and the service contacted him to say 3 3 there anything anyone wants to say about that wasn't possible, and he had to wait 4 infection and diagnosis of hepatitis C? 4 a couple of weeks to find out why, and the why 5 PAM: 5 was because he had hepatitis C. He was No. JACKIE: No. 6 6 absolutely devastated. This was a man who had 7 KAY: No. 7 already, once he'd got diagnosed, as a result 8 MS SCOTT: The next theme you pick up, and you've 8 significantly in terms of his relationship with 9 9 already referenced this, is infection of blood his children, because he was so scared of 10 10 donors, Kay, do you want to start us off on that infecting them, and then at the same time, his 11 theme? 11 mind was around the guilt, and, you know, it's 12 KAY: Yeah, several people were concerned that 12 very easy to say well it's not his to have. He they'd been blood donors and that they'd 13 13 felt guilty guilt. He still feels guilty that 14 actually passed on the infection without their 14 other people, throughout that 15-year period, 15 knowledge, and we've kind of discussed this and 15 would have been infected and he was doing a good 16 put this under this theme in of good people 16 thing having had his life saved. 17 doing good things. 17 I think it brings up the issue that we talk 18 You know, one gentleman that I met was 18 about later on, which is in the absence of any 19 17 years old and had been involved in a road 19 psychological services, specialist psychological 20 traffic collision, and as a consequence of that, 20 services with people who can get the support to 21 has had life-saving treatment in the hospital, 21 work through that and feeling guilty, and to 22 22 and was so so grateful that that had been done, kind of work through that that's not something 23 that he went on, like others, to be blood 23 that they have to bear, he today lives with that 24 24 donors. tremendous guilt, and it continues to impact on 25 25 15 years later, he was waiting for his next all his relationships in the here and now, and 39 40 1 he still has hepatitis C. So yeah. That's, you 1 of people she may have infected during that 2 2 know ... Pam? 3 PAM: A couple of examples from me ... we met 3 Another example was a woman who had had an 4 a woman who had blood transfusion when she was 4 ectopic pregnancy and she was a very rare blood 5 5 group so she was encouraged and obviously wanted 14, and she started to be a blood donor when her 6 age permitted, and she then discovered in her 6 to give blood which she did, and continued to do 7 7 30s that she had hepatitis C. This lady was that for 20 years and during that time had 8 pretty tremendous in so many ways. She now 8 possibly infected people and lived with the 9 9 worked in a factory with -- actually depot, with guilt of that also. 10 lots of men. In fact, she's the only woman 10 (Off the record comments from Sir Brian) 11 working in the depot and very, very keen to keep 11 (11.30 am) 12 her cup separate and all her individual items 12 (A short break) 13 separate because of her fear of cross-infection. 13 (12.00 pm) SIR BRIAN LANGSTAFF: Yes. 14 But the blood donor van, as we call it, comes 14 15 MS SCOTT: So we were, Jackie, I was asking you just round, the mobile van, to the factory to take 15 16 blood donor, and all the men go out to give 16 before we broke, to move on to the next theme, 17 blood, and she can't and doesn't. They don't 17 which was other infections. 18 know why she can't and doesn't, and they call 18 JACKIE: Yes, we saw a small number of people who, 19 her names like scary cat, et cetera, because she 19 around about ten to 12 years ago, in their 20 wouldn't give blood. 20 memory, were advised that they might have been 21 21 exposed to variant CJD through their treatment. Clearly she can't share with them why she 22 22 And whilst they felt some reassurance that this can't give blood, and the -- listening to her 23 23 account, because she was such a good woman who was being properly investigated, the manner in 24 given blood for years, but then as Kay was 24 which they had been informed of that was really 25 explaining, felt so so guilty about the number 25 quite brutal.

42 1 They all spoke about receiving a letter 1 recalled any of the risks of transfusion being 2 completely out of the blue, and that wasn't --2 discussed with them by professionals at the 3 3 there was no phone call, there was nothing via time. And it was just accepted with those who 4 their GP. And the letter said, in a very formal 4 had haemophilia and thalassaemia that they would 5 5 receive these products, and almost like, "This manner: 6 "We understand you have received blood or 6 is a wonder drug", and there was this kind of, 7 blood products from someone who has been 7 you know, we're really grateful for this, but 8 diagnosed with variant CJD. Would you like 8 none of those risks and the issues around 9 a test?" 9 consent and any discussions were recalled or 10 And they, for some of them, this was the 10 very few were recalled. Particularly, I wanted 11 period when they were not only managing the 11 to make reference to a lady that I met who'd 12 symptoms of the disease but they were in that 12 been informed by her consultant that she'd 13 period where they were having interferon, 13 sought out herself a second opinion, because the 14 ribavirin treatment as well, and it was just the 14 treatment that she was being given in another 15 last straw for some of them, and one of them 15 hospital was interferon only, because she had 16 said, you know, "What else? What else is going 16 genotype 1 hepatitis C, and as a consequence of to be thrown at me?" in that respect. 17 17 that, she was being given the treatment, and she 18 So that was a relatively small number of 18 was really struggling with the side effects, and 19 people but a big impact for them at that time. 19 it was getting to the stage in terms of that it 20 MS SCOTT: Kay, can I ask you, then, to move on to 20 was impacting on every part of her life and her 21 the next theme you pick up in the report which 21 family's life, and she wanted to come off this 22 22 is consent. or at least know some more information. 23 23 And she felt real pressure from the first KAY: Okay, so very few people could recall being 24 24 asked for consent about the use of blood professional to continue, and this went on for 25 25 products or a blood transfusion, and nobody quite a long time, until she eventually said no, 43 44 1 enough's enough, and she sought a second 1 consider the implications, but none of them --2 2 opinion. She went to a completely different it wasn't explained to them about the possible 3 hospital, and had a very different experience, 3 risks of having transfusions. One person's 4 but what she also learnt from the second 4 parents were telephoned in the middle of the 5 consultant was "You are on a trial" and she had 5 night to give consent, the person was 16, and 6 no idea that she'd been on a trial. She'd had 6 she'd had a haemorrhage after an operation. And 7 7 but they were told they were asked to give no discussion about it, she'd not consented to 8 8 consent verbally and they were told it was in 9 9 the context of life-saving procedure. And this is in contrast to another gentleman 10 that I met in a different area of the country 10 And they and other people reflected on the 11 who also had genotype 1 hepatitis C, who'd had 11 guilt that that brought for them, that they had 12 a really long conversation about interferon only consented for their child to have that 12 13 treatment, and its success. And actually, he'd 13 transfusion. 14 made a decision, because it was so low at the 14 Lots of people said to us there was this 15 time in terms of, you know, what could happen, 15 terrible irony of having their life saved on the 16 he'd not -- anyway, he'd waited until 2015, 16 one hand, but being given something which would 17 until the new drugs became available, and that's 17 ultimately limit their life on the other hand, 18 when he made a different decision. But it 18 and that was very difficult to deal with. 19 brings out this real issue of consent around 19 There was a specific issue that someone 20 risk, and consent around trials. 20 explained to me, and this was a person who, in 21 21 a relatively new relationship, was informed by And I was going to hand over to Jackie, then 22 22 her partner that he had HIV infection, and it to talk about another issue relating to consent. 23 JACKIE: Okay. When -- lots of people who had 23 was agreed that she would go to his consultant 24 transfusions in an emergency said obviously that 24 to have some tests, which she did. But she was 25 they weren't in a position themselves to 25 also tested for hepatitis C. She didn't know

45 46 1 that the partner had hepatitis C, and so she 1 impairment was often described as "brain fog" to 2 wasn't asked for consent or told that she was 2 us, and that led to a debilitating level of 3 3 being tested for hepatitis C. confusion and inability to focus that was 4 When she took this up later with the 4 sometimes combined with associated anxiety and 5 5 consultant, she was told that she couldn't be depression, and quite a few people spoke to us 6 told that she was being tested, because it would 6 about losing their way on the way home, not 7 have compromised the confidentiality of the 7 being able to remember journeys. Lots of people 8 other person, and she felt really strongly that 8 were really anxious about driving, for example, 9 9 her rights, her welfare, had been overridden by because of the impact of that upon them. 10 10 the confidentiality of someone else. People have also experienced mental health 11 So the next section is we want to talk about 11 problems to a significant effect, which really 12 people living with the infection, and we're very 12 impacted on everyday life, their relationships 13 much minded that this audience can tell us a lot 13 as well. People described the later stages of 14 more than we could ever explain to you. I'm 14 the disease, again physical issues, where the 15 going to speak to the accounts that were given 15 liver and organs were affected. Living with the 16 to us about impact on individuals of that. 16 worry about varices and the possibility of those 17 People talked about the impact of 17 bursting there -- obviously enlarged veins. 18 hepatitis C particularly in vivid detail, and 18 People having to go through really painful 19 19 the really common issues were chronic and invasive tests, particularly in the early 20 debilitating fatigue and how that impacted on 20 days when liver biopsies were very invasive and 21 their lives, often lasting for years, as we've 21 involved. Without going into too much detail, 22 said earlier, without diagnosis. 22 long needles and breaking through the skin. 23 Other symptoms were severe aches, 23 One witness said to me, "As a result of my 24 24 gastrointestinal problems, skin changes and infection and the worry around it, I've suffered 25 25 sweating, these are physical issues. Cognitive mental strain and stress to the point of 47 48 1 thoughts of suicide, out of my fear of having 1 children of that family. 2 a nasty death." 2 So the different elements that were 3 So it wasn't just about where people are at 3 happening there together with, within some 4 the moment, or were at the moment, in the course 4 families, you've got more than one infected 5 of their disease, but that kind of anticipation 5 person, particularly with families where there 6 about where this would lead to. And the point 6 was haemophilia and we got brothers, sons, 7 7 we would want to make was, even though we've fathers, et cetera. So you'd got a number of 8 seen -- the majority of people we've seen have 8 layers of different tensions and pressures for 9 9 now been cleared of the illness, lots of them those families. And the impact on keeping the 10 are still experiencing some of the symptoms of 10 secret, and meeting a number of people who 11 the illness itself, the impact of those, and of 11 actually didn't want to have discussions within 12 the treatment which we will come to speak about the family because they didn't want to burden 12 13 13 the rest of the family with what they felt was 14 PAM: I want to briefly speak about the impact on 14 their condition. So to have your children 15 the family and the relationships, and there's 15 having to understand that condition, they felt 16 a number of strands to this element. The strand 16 was burdensome, so kept a lot of that to 17 of the pressure in a family of keeping a secret, 17 themselves, which created stress, anxiety, and 18 the stigma, if the outside world was to realise 18 often depression in the way that Jackie has 19 what was happening by way of the infections. 19 relayed it. 20 But also, the actual daily routines in families 20 So we'd got families who certainly had a 21 21 feeling of secrecy and an underlying, almost which were creating lots of stress: the need or 22 22 perceived need to keep separate towels, to keep depression, but certainly an anxiety within the 23 23 separate crockery, to make sure that there was families which was palpable often. 24 no bodily fluids exchanged between the infected 24 I think Kay was going to talk to us about 25 person in the family, on the adult, and the 25 a specific case that she'd been in contact with.

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that she and any children she had could have

been infected too by having a relationship with

50 1 KAY: One of the families who I met had a child 1 that this holding the feelings down, sometimes 2 who had been infected with HIV, and I think 2 even in the most close relationships because 3 3 probably to explain the impact on family and actually they were just surviving the moment, 4 relationships, the best way is to use her words, 4 trying to put one foot in front of the other, 5 and what she says is "We have nothing else to 5 day by day. So yeah, that's them and I'll just 6 compare this to. Our child was infected and it 6 hand back to --7 took the joy out of our lives. Psychologically 7 JACKIE: To me. 8 we became different people. We're cut off from 8 The business of keeping secrets was a real 9 9 others, and the depression we carry has at times strain for people, we found. And a few people 10 almost torn us apart. In the darkest days, I 10 said to us they had to learn new techniques of 11 even chose the music for our son's funeral." 11 when they were in company with people and to be 12 And although this couple feel extremely 12 able to manage the kinds of things that they 13 lucky that their son is still alive, what we 13 talked about in case the conversation went down 14 talked about was the impact this has had on them 14 a particular route, and they found themselves 15 as couple, and within their family relationships 15 saying something which they then had to 16 outside of that, and with their other son. And 16 extricate themselves from. 17 one of the things that we commonly found was 17 So not at all expecting in their lives that 18 that we would sit in maybe the front room of 18 they would have to do that. And one chap 19 somebody, we'd be talking to two people, and 19 I spoke to said, "I'd never lied to anybody in 20 they'd be sharing their thoughts, feelings, and 20 my whole life before", and that was a really 21 exactly what had been going on for number of 21 hard experience about not being able to tell the 22 years, and one part of the family member would 22 truth to his workmates about his condition, and 23 say to the other, "I never knew you felt like 23 quite a few people said that they had other 24 24 that. I never knew that had gone on for you." chronic conditions when they were asked about 25 25 And that was a common thing that came about, their symptoms as well. 51 52 1 Going back to the issue about personal 1 me." 2 2 relationships, we did see a few people who sadly And that was terribly, terribly sad for us 3 found that their closest personal relationships 3 to see that. 4 were affected. One man that I spoke to who had 4 PAM: I think probably people in this room will 5 lost his brother to HIV, when he was -- when he 5 all remember the medical advice in the 1980s for 6 was 20 and his brother was 23, "Growing up, he 6 men co-infected with HIV, and hepatitis C, which 7 7 was my best friend. We were inseparable in the was to abstain from any sexual activity which 8 school holidays but his HIV diagnosis drove 8 involved exchange of bodily fluids. And at 9 9 a wedge in our relationship. It was something least two of the men interviewed remained 10 that we never discussed properly and we were 10 celibate throughout their twenties after 11 never close after that, and that's one of my 11 receiving this advice, and I say "at least" 12 biggest regrets and I feel guilt to this day". because clearly this was a sensitive and 12 13 We saw people who made the decision, as Pam 13 difficult subject for any of us to be talking about with the people we interviewed. So it was 14 has alluded to earlier, about just not taking 14 15 the risk of having an intimate relationship, of 15 quite difficult to actually get the extent and 16 starting a relationship, with a partner. 16 understanding of how this had affected people in 17 I have to say we met some wonderful people 17 all kind of parts of their relationships. 18 who would have made wonderful partners, 18 So the two men who talked about being 19 wonderful parents, and they had had choice --19 celibate, one did marry later, but the other 20 they felt they had that choice taken away from 20 never felt able to have a relationship for fear 21 them. And I think one man really summed that up 21 of infecting a partner. 22 22 for me and said, "It's hard to go on a date with Similarly, the advice on sexual activity was 23 23 a woman and tell her you've been infected, and not clear for people infected with hepatitis C,

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and everyone who we interviewed described

significant stress because of this, as we've

53 1 said. Even when they were in very close, loving 1 though there'd been a recovery from hepatitis C, 2 relationships, it was still an area that was 2 the future was limited in so many ways, with 3 3 very difficult, and probably one of the most a terrible, terrible sadness ... 4 difficult areas for us to explore and understand 4 KAY: I just wanted to follow that with looking at 5 5 those that were -- the impact in communities. entirely, other than we knew that the amount of 6 stress that this certainly brought with it. 6 So one of the families that I met, and those 7 We'd met a number of couples who spoke with 7 with underlying conditions, with thalassaemia 8 immense sadness about limiting their families 8 and haemophilia, developed friendships with 9 9 because of hepatitis C ... others at some of the places that they were 10 10 Obviously on a number of occasions it meant going to have their treatments and their 11 that until they'd had their infection cleared, 11 transfusions, and one of the real impactive 12 or noticed that the infection was cleared, some 12 things was a gentleman who said to me "We were 13 were older. Fertility had diminished in some 13 like a little family. We were in and out all 14 cases. Others just felt they could not risk the 14 the time. And it was hard seeing them die." 15 infection returning if they had children. And 15 And how that impacted on him, and his life, and 16 emotionally being spent, in the sense of even 16 still does, today. 17 beginning to think about the responsibility of JACKIE: I spoke to one woman who, unbeknown to her, 17 18 children. 18 was infected during a transfusion after the 19 19 birth of her first child, and then had some We spoke with a number of women who had 20 terrible, terrible sadness as their lives had 20 trouble conceiving later on, when they wanted to 21 been about having children at the appropriate 21 extend their family. And she was referred for 22 point, and that had now been something that was 22 IVF, and obviously part of the that service was 23 lost to them. 23 that she had to have some blood tests, and she 24 24 So we certainly -- I think those were some received a telephone call from the hospital 25 25 which said, "You are no longer suitable for IVF of the very, very sad interviews, that even 55 56 1 because you have hepatitis C." 1 we've said, people lived with an enduring sense 2 2 And that was how she then found out about of feeling blamed and feeling shamed about the 3 that. She did then conceive naturally, and she 3 infection. Which was really compounded by the 4 said that there was real trauma, then, 4 national mood: lots of people said the whole 5 surrounding her second child's birth, which was 5 thing was mixed up about the national mood in 6 highly, highly medicalised. When he was born, 6 the eighties, in particularly around HIV and 7 7 he had to have his stomach emptied via suction AIDS and the judgements that were placed on 8 to ensure that no blood had been ingested and 8 people in respect of that, and a sense of moral 9 9 she said the midwives took him to the other side panic, really, that they felt very frightened 10 of the room to do that so she wasn't able to 10 about. 11 hold him for guite some time. Eventually, they 11 More than a couple of people talked to us 12 were both moved from the labour suite and placed about being called junkies in the street and in 12 13 her in a side room away from other patients, so 13 other wider friendships. Some people didn't 14 that she couldn't mix with other mothers, and 14 tell their children at all about their 15 she said that was a highly traumatic period of 15 condition, as Pam was saying earlier, and that 16 the first hours and days with her second child, 16 was because they didn't want their children to 17 and that really impacted on her as a parent as 17 be bullied at school. They were really afraid 18 she went forward. 18 of that ... MS SCOTT: Jackie, I'm going to ask you now to turn 19 I saw one woman who was really acutely 19 20 to the next theme you pick up in your report 20 anxious about anyone finding out that she was 21 which you've alluded to already, quite a lot, 21 infected. And the receptionist at her 22 22 child's -- at her GP surgery was also a parent and that's stigma and shame. 23 23 JACKIE: Mm-hm, this was a huge issue for absolutely at her child's school, and she went through such 24 everybody that we saw. And you'll have heard 24 agonies every day wondering whether this 25 that it's mapped through the other things that 25 receptionist might have shared, inadvertently or

	57		58
1	in any other way, information about her		had haemophilia, therefore he might have HIV.
2	infection. And this led to a really crippling		d society at that time and in the media, there
3	anxiety. So the family felt obliged to move		s some high profile kind of footage being
4	house, and they moved 25 miles away, away from		eased about HIV that was creating this fear
5	all of their support networks. And then,		d scaring the community, and that was leading
6	four years later, she had a new neighbour who		a real lot of prejudice and discrimination in
7	was a GP at a local surgery and again, this		ciety. So not only did we have these internal
8	triggered her anxiety, and the family had to		ues of managing family relationships,
9	move again.		naging yourself, you've then got the societal
10	And the issue there was that there were		pact of the prejudice that was out there, the
11	additional burdens for people about dealing with		r that was out there, and how that was
12	this situation about having to move themselves		ndled and managed at the time, was adding to
13	away from the support networks which were		psychological stress of our families.
14	helping them through it.		OTT: Pam, did you want to add anything?
15	<b>KAY</b> : I just wanted to touch on the issue that	15 <b>PAM</b> :	Just bringing us to present day, really,
16	we've mentioned there about society's response		it we know that people felt there was an
17	to the blood infection through the time, and	_	orance around hepatitis C, but a need to
18	specifically in relation to prejudice and		are some of their status with others, as
19	discrimination, and one family I met talked		leed to protect others. But what they were
20	about in the eighties, having a sign in their		ping was, from the Blood Inquiry, that there
21	local area displaying a sign that said, "No		s much more of an understanding within our
22	homosexuals or haemophiliacs", and what people		mmunities about infected blood and how it was
23	were doing with that led to another gentleman		used, and that publicity from the Inquiry
24	who was haemophiliac having to have his own		uld raise awareness and help to quash these
25	glass in his local pub because people knew that	25 ill-ir	nformed myths that were around. So they
	59		60
1	brought us into present day as to what they	1 free	edoms that they hadn't previously
2	hoped from, from the Inquiry.		perienced. Obviously in retrospect they all
3	MS SCOTT: Pam, can I ask you then to pick up on the		mmented on the Catch-22 of the long-term
4	next theme in your report, which is an impact on		olications of the resulting infections. So we
5	education and work.		d this mixed picture of, yes, "We enjoyed
6	PAM: Yes, certainly.		ng in these schools, and they gave us the
7	We've heard that we've met with a number of		edom that we hadn't otherwise had", and there
8	people with haemophilia, and we know that in the		s a good understanding of their medical needs,
9	seventies and eighties, it certainly wasn't		then, as I say, we have the implications of
10	uncommon for children with haemophilia to be		e clotting agents.
11	educated at what we know as special schools	11	l also met with two women with thalassaemia
12	during that time for pupils with additional		o actually had very different school
13	health needs.		periences, and that may have been due to the
14	And I interviewed two men educated in one of		t that there was a 15-year age difference
15	these schools who actually had very good		tween them. The older woman found that
16			
	memories of the school at that time, and they		lassaemia led to fear and exclusion by other
17 10	were supported in managing their haemophilia.		Idren. She required frequent blood
18 10	And injecting themselves to avoid hospital, and		nsfusions and was absent from school for
19	they explained to me how, prior to being able to		ignificant amount of time, and she doesn't
$\alpha$	inject themselves, there were frequent visits to		all any additional support during those
	hospital, and that clearly interfered with their		ars. I actually interviewed this woman while
21	and the state of t	22 she	e was having a blood transfusion, I sat with
21 22	education and their ability to spend time in and		
21 22 23	around other young people.	23 her	r in hospital during that time, and her whole
20 21 22 23 24 25		23 her 24 life	

1 bright woman, with lots of energy, but her 1 who had had to give up on their education, for 2 schooling had been entirely wiped out by her 2 example I saw one family where a young woman had 3 3 need for blood transfusions and then, later on, been to university and had to give it up after 4 an awful lot of treatment. 4 a year because her father was in the end stages 5 The younger woman with thalassaemia was well 5 of his illness and she couldn't bear to be away. 6 supported at school and college and her 6 She said that every time she went back to 7 experience at work was not as positive. One of 7 university she feared if she would see him 8 her employers required her to make up the hours 8 again, and so she gave that up, and wasn't able 9 9 missed for her transfusions, resulting in long to pick up with her learning for several years 10 10 days, and further exhaustion, which we knew she after that. 11 experienced through the hepatitis C. Clearly it 11 But we also saw people who gave up on 12 was not sustainable to have to make up her time 12 education when they were infected. Either 13 at work when she'd been to have treatment and 13 because of the pressure of treatment at the 14 blood transfusions. The days just weren't long 14 time -- I saw one man who was trying to take 15 enough for her to do that, it felt an entirely 15 professional exams whilst he was having 16 unreasonable expectation of her. So she moved 16 interferon and ribavirin treatment, and he said 17 employers for those reasons. 17 one day he'd just slept in for his exam because 18 Fortunately, her current employer is much 18 he was so exhausted, and there was no quarter 19 more supportive and flexible and she's able to 19 given about that. He just had to miss out on 20 lead a more regular life with this level of 20 his exam. 21 support. But for the two of them, it was very, 21 People were experiencing mental health 22 22 very restricting, or restrictive. difficulties, and didn't feel able to continue 23 MS SCOTT: Jackie, did you want to add anything 23 with their studies. And a couple of young men 24 24 about employment? who just said they thought it wasn't worth it 25 25 JACKIE: Yes. We saw quite a few examples of people because they thought their life expectancy was 63 64 1 going to be so limited, and that they didn't 1 nature of their work or working for five days 2 2 want to invest their time in education. a week was really just too much for them. 3 3 And people then spoke about how there was On the other hand, several people spoke to 4 us about their absolute determination to keep on 4 a loss to society as well, because of that. So 5 working in spite of their debilitating 5 some of the examples that we saw were about, you 6 condition. And sometimes this was about wanting 6 know, a woman who was a nursery nurse who had to 7 7 to contribute to society, and not giving in, retire at 49, a nurse who had to retire at 52, 8 feeling that they didn't want to give in to the 8 a teacher who had to retire early because her 9 9 illness and allow it to take yet more away from mental health problems, as a result of her son's 10 them. But for others, it was a financial issue 10 condition, were just too much for her to feel 11 and they just couldn't afford not to work, and 11 that she could carry on with that. 12

so they had to carry on. And I saw a couple of people who said they felt they had to drag themselves to work in their sixties until such a time as they reached their statutory pension age, because they just couldn't afford not to do that. And they spoke about the -- you know, the impact of trying to work with carrying those symptoms of the illness, and people spoke about a life consisting of working and sleeping, really, for years, just trying to keep going. People having joint replacements, that kind of

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thing.

Several people had to reduce their working hours, or change careers, actually, because the MS SCOTT: So now I'm going move on to the next section, which is treatment, care and support.

Can I ask you, Jackie, to start with some of the experiences you heard about the attitude of healthcare professionals to those that you spoke to?

JACKIE: Yes.

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Our view is that anybody who's experienced a serious illness, or any illness, knows the importance of -- the manner in which a person is treated is often as important as the treatment itself. And we heard a range of experiences from people, from sensitive and expert care, which they really appreciated, to brutally

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66 1 delivered information, inappropriate questions, 1 a flexible and practical response to the 2 and frankly poor quality care. 2 challenges they faced, and probably came the 3 3 Lots of people described their consultants closest to any kind of therapeutic help for people in the manner in which they conducted 4 as lovely. They shared examples of medical 4 5 5 nursing staff absolutely going the extra mile to their work. 6 support patients with home visits, open-door 6 One family described a really positive 7 access. Some of the consultants attended the 7 experience of care for their uncle who had 8 funerals of people who had died. One woman was 8 haemophilia. And this was a man who had been 9 widowed when her husband died, said that she 9 born in the 1920s, so he'd undertaken some 10 didn't know how she would have survived without 10 extreme treatment in his youth when people were 11 the support of the consultant, and in the course 11 finding a way, and he was under the care of the 12 of the work that we did, we were able to trace 12 same two consultants and a very dedicated 13 where the consultant had got to in the country, 13 nursing team throughout his life. And the 14 which was nice for her because he'd moved on. 14 team -- the family said the team were like 15 Quite a few people spoke to us about, 15 a second family to them, and their lives 16 particularly those with haemophilia, who said 16 revolved around visiting the unit, and when the 17 that their own consultants felt a real anger 17 family went back to the hospital, as part of 18 about what had happened to them and they were 18 their work to support this inquiry, the 19 active campaigners around infected blood and 19 receptionist remembered them from 30 years 20 really working hard to find new treatments to 20 earlier, which was an indication of that real 21 tackle hepatitis C in particular. 21 sense of family that people had. 22 22 One thing that we wanted to stress was And they felt, in terms of their uncle, that 23 almost exclusively, people spoke very, very 23 given the extent of his haemophilia, they did an 24 positively about the specialist nurses that --24 amazing job keeping him alive into his sixties. 25 25 the specialist hepatitis C nurses who provided And he actually died when he was 65, as a result 67 68 1 of hepatitis C, unfortunately, and at that stage 1 certainly his treatment, he said, couldn't be 2 2 he was the oldest haemophiliac in his home city. better 3 So they had some real mixed feelings about the 3 He of course was very aware that the blood 4 clinicians had really worked hard over many 4 transfusions which were there helping to keep 5 years to keep him well and to keep his 5 him alive in the first instance, became 6 longevity, but it was such a shame that after 6 a roller-coaster of events that, following that 7 7 all that excellent care, that he passed away as heightened feeling that he was being treated 8 a result of the infection. 8 successfully, then came the news about this 9 9 Pam has got some other examples. hepatitis C and needing to continue to have 10 PAM: Yeah. I met with a man who had a very 10 different treatment. 11 positive account and experience of his 11 Fortunately, the day that I visited him, 12 treatment. He had developed Hodgkins lymphoma he'd just had the all-clear from his hepatitis C 12 13 in his late teens and had been treated for that 13 treatment and was in a very good place. So we had a good outcome. And certainly we've been in 14 very successfully, and enjoyed good 14 15 relationships with his consultants, but he was 15 contact around our presentation today and he 16 contacted by his consultant around '95, '96. 16 continues in good health. So, as I say, a very 17 The consultant was extremely apologetic that the 17 positive experience which was very heartening to 18 infected blood had been given at the hospital 18 19 where the consultant was working. 19 I think Kay has got an example too. 20 And the man I was interviewing was extremely 20 KAY: Yeah, I just wanted to add to what Jackie 21 keen to emphasise that the specialist care he 21 had mentioned about specialist nurses and some 22 had received for his Hodgkins lymphoma and 22 of those doctors and nurses that had been there, 23 23 hepatitis had been excellent. And he said when and were real positive experiences. 24 he visits his GP, he welcomes him like a VIP, he 24 In contrast to that, where specialist 25 is always made to feel very special and 25 facilities were not available, people described

69 70 1 being treated in clinics alongside drug and 1 there, they were signposted and got some help 2 alcohol users, or with cancer patients, and 2 and support and managed to keep their home, and 3 3 I think that was mentioned earlier on. If they obviously their children live with them there as 4 lived in an area with low incidence of 4 5 hepatitis C, this also limited their ability to 5 So it made such a difference, and that is 6 establish networks and gain support from other 6 a service that is out there in one of the areas 7 people who were also affected. 7 that we came across. 8 However, in contrast to that, in the 8 PAM: In stark contrast to my last example, I've 9 families that we met, I met one specialist 9 a pretty harrowing example from another family, 10 social worker who is working with the 10 and one man recounted that his father had 11 haemophiliac community in that area, and it is 11 leukaemia and received stem cell treatment 12 a designated post, a specialist social worker, 12 followed by blood transfusions. He was at this 13 in relation to everything in and around 13 point infected with hepatitis B, which was noted 14 psychological support, financial support, 14 as the primary cause of death on the death 15 general advice, and to use their words they 15 certificate, and I think all three of us have 16 said, "We'd not be where we are without her 16 seen a number of death certificates throughout 17 today. She's helped us deal with financial and 17 this, at our time working in the Inquiry. 18 emotional matters. We rely on her and everyone 18 His father had been told by the consultant 19 19 that an error had been made and the wrong blood deserves this support." 20 And where services didn't have that, others 20 taken out of the wrong fridge. This had 21 spoke of feeling isolated, and abandoned, and 21 affected eight people, seven of whom had died. 22 22 that specialist service made so much difference The father was told he may be all right, but in 23 for a family who were going through bankruptcy 23 fact he died three months later. 24 24 because their business had to fold, and Again, that sounds like a pretty strange 25 25 actually, through the specialist knowledge out story, but they were able to show me a statement 71 72 1 that the father had given to his solicitor at 1 really sunk in what has happened. Now that 2 2 the point that he was informed about this, and I have hepatitis B, I cannot be cured, and there 3 they'd still got the statement, which didn't 3 is a possibility that it will be passed on to 4 ever go anywhere in legal terms, but was there 4 others who come into contact with my blood." 5 as a record of this meeting with this solicitor. 5 He died very shortly after this, and I'll 6 Following on from that, the family were 6 come on later to talk about the circumstances 7 7 distraught about their father's treatment as he under which the family heard about his death. 8 spent his last weeks in an isolation ward at 8 Handing over to Jackie. 9 a psychiatric hospital. And this was a man who 9 JACKIE: Just to pick up a couple of things from 10 was a businessman. He'd fallen ill only a few 10 what Pam said about death certificates. I had 11 months prior to this, because he was having his 11 two differing experiences. The young man 12 successful treatment for leukaemia, and then, as that I mentioned earlier who'd died from HIV, it 12 13 I say, became very ill very quickly. So to have 13 wasn't mentioned on his death certificate. His 14 his last days in a psychiatric hospital in an 14 death certificate said that he'd died from liver 15 isolation ward, at this was due to the lack of 15 failure, and the explanation to the family was 16 provision to treat his hepatitis B in the main 16 that the medics had done them a favour by not 17 hospital, and the fear from the hospital workers 17 mentioning the HIV in that respect. 18 that he was highly infectious, and the father 18 With another family that I worked with, their doctors were very angry that their 19 wrote this account when he was told about the 19 20 infection which as I say was formulated into his 20 patient, who had haemophilia, had been infected 21 21 with hepatitis C, and they were determined to statement and I'll read some of the words which 22 22 make sure that that was noted, and on the death are in this statement and they were able to show 23 23 certificate itself, the doctor had written 24 "I was told today by the consultant that the 24 diagonally right across it in large letters 25 blood cells were contaminated. It has not 25 "hepatitis C", and the woman who I worked with

73 74 1 said that when the registrar opened it she was 1 that essential tests and equipment weren't 2 quite taken aback, and the message was that the 2 always available on the NHS for people, and 3 3 clinicians really wanted people to know the people were expected to fund them privately if 4 cause of that. 4 they wanted to have these newly developing 5 5 facilities. And in 2005, one patient was told In another example that I have, I saw a man 6 who had had successful treatment to clear his 6 that a fibroscan would determine the progress of 7 hepatitis C, and he and his family felt they 7 the disease, but there were no facilities in 8 wanted to move to a city, a new city, for a new 8 this country and she had to travel to Paris in 9 9 start. So his treatment for haemophilia June of 2005 at the family's expense. The 10 10 transferred to a new hospital. And on his first following year, a fibroscan machine was 11 appointment the doctor that he saw said to him 11 installed at a hospital 74 miles away from where 12 "Oh, I see you've got hepatitis C", and the chap 12 she was, and she had to travel there for scans 13 said it was a terrible shock to him, as he 13 for 7 years before she could have them more 14 thought the infection had cleared, and he was 14 15 told to come back six months later for 15 Then in June of 2008 a consultant advised 16 a follow-up appointment, and when he went back 16 that, due to the level of cirrhosis, she needed 17 after six months he was told that this was 17 to have an enhanced liver fibrosis test. Again, 18 actually a clerical error and what should have 18 this was not available on the NHS, and she had 19 been written on his record was that he was 19 to pay £99 to have this carried out by a private 20 positive for hepatitis C antibodies and had been 20 contractor in an NHS hospital. 21 wrongly copied as positive for hepatitis C. So 21 And just that -- we wanted to share with you 22 he spent six more months thinking that, and he 22 that the words that people most frequently use 23 said that really put him through months of hell. 23 when discussing access to treatment were "leper" 24 24 Another issue that came up for us around or "pariah". That is how they were made to 25 25 treatment, particularly in the earlier days, is feel. And there were many, many experiences of 75 76 1 people being denied treatment or isolated from 1 In contrast, there was an example of one 2 2 other patients because of hepatitis C, and we woman's dental nurse commenting loudly -- in the 3 saw quite a few people who had difficulties with 3 infection, but the dentist was challenged -- the 4 dentists, for example, and Pam has an example 4 nurse was challenged, I should say. So there 5 there to share -- oh sorry, it's Kay. It's 5 was some good evidence within professional 6 Kay's, I'm sorry. 6 environments where people were not behaving in 7 7 KAY: Just moving on, we're talking about the an okay manner, but there was also some evidence 8 dentistry. One of the families that I met had 8 of other professionals challenging them and that 9 9 had such poor treatment in relation to really making a difference for the individual, 10 dentistry, where the procedure had to be 10 in terms of that prejudice and that 11 undertaken in the hospital, the hazardous 11 discrimination. 12 stickers were out, it was at the end of a day, MS SCOTT: We've come to the end of that theme and 12 13 the attitude of staff was not good for him at 13 we're going to move on to another one, for treatment of hepatitis C, but I notice the time. 14 all, that it left such a significant impact on 14 15 SIR BRIAN LANGSTAFF: So this would be a convenient his life that, for the rest of his life, he's 15 16 never gone -- he's still alive today, he won't 16 moment to have our lunch break, would it? 17 go to the dentist because the experience was so MS SCOTT: Indeed. 17 18 poor that this has led to -- when he has needed 18 SIR BRIAN LANGSTAFF: Can I thank you very much so 19 a couple of extractions, he's done that himself. 19 far, and I hope you don't mind having to come 20 And actually, you know, if you think of that 20 back after lunch. It will be two o'clock. But 21 21 what you have so far told us, I think, has been in context to the pain that you must have to go 22 22 really valuable and worthwhile listening to. So through in terms of extracting your own teeth, 23 you would have to have suffered so much of 23 thank you so far, and I look forward to seeing 24 a psychological impact to do that, that it's 24 you back at two o'clock. 25 astounding. 25 Two o'clock

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1	(12.58 pm)	1	the treatment before the other, whereas others
2	(Luncheon Adjournment)	2	felt that they entered it quite naively. And
3	(2.05 pm)	3	I think all three of us heard the phrase that we
4	SIR BRIAN LANGSTAFF: Yes, you're moving on to the	4	had heard from a number of people that said,
5	next topic.	5	"This was the worst year of my life". People
6	MS SCOTT: Yes.	6	certainly articulated suicidal thoughts and
7	Pam, you were going to start us off on	7	people seriously had contemplated taking their
8	treatment for hepatitis C.	8	own lives because they could not see an end to
9	PAM: Yes, thank you.	9	how they felt
10	To take people right back to the beginning,	10	They described a very, very debilitating
11	when we explained how many people we'd seen with	11	illness, extreme tiredness, staying in bed for
12	various disorders and illnesses, and we actually	12	days and days and days, not coming downstairs,
13	met with 50 people who were infected with	13	literally for weeks, to be with their family.
14	hepatitis C, and I think it's fair to say on	14	Hair loss, significant weight loss, and also
15	behalf of all three of us, that it was truly	15	a profound impact, not surprisingly, on their
16	shocking to hear for the first time just the	16	own emotional and mental health.
17	experience of these people who had been treated	17	Many of the effects of the treatments have
18	with interferon and ribavirin. They described	18	endured and have become lifelong conditions.
19	it as gruelling, similar to chemotherapy, which	19	I need to talk about a young woman with
20	lasted months and, in some cases, years.	20	thalassaemia who had been engaged to be married
21	A small number of these people said they	21	when she started the treatment. She was in her
22	were given clear information to help them	22	early twenties then. Until that point she'd had
23	prepare for the impact of the treatment, and	23	a life where she had been very successful, she
24	a good example here was a couple who had both	24	had managed her thalassaemia well with a very
25	been treated and able to plan that so one had	25	supportive family.
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1	Then learning about her hepatitis C and	1	And the effects of the virus and the
2	starting with the treatment absolutely changed	2	treatment have left her with enduring severe
3	her life, to the extent that she certainly	3	depression and anxiety. She describes becoming
4	suffers with a great deal of depression,	4	mentally exhausted very quickly and uses the
5	anxiety, loss of confidence, and she talks about	5	word "overwhelmed" a lot in terms of how she
6	not leaving her parents' home for a year.	6	feels.
7	Her wedding that had been planned had been	7	She described the overall result of the
8	cancelled, was cancelled, the relationship	8	infection and treatment as "an existence of
9	ended. She was unable to work or consider any	9	survival and coping with little contribution or
10	career prospects.	10	fulfilment".
11	The anticipated future of leading a life	11	I met one family whose relative had an
12	with a partner and a child, and a professional	12	uncommon, we think, reaction from the treatment,
13	job that she had, were all lost. Sadly, she's	13	which was a very significant weight gain, and
14	now got cirrhosis of the liver and is deeply	14	the implications of that. This lady had been an
15	pessimistic about her life expectancy. A very,	15	average weight, looked after herself very well,
16	very sad story. We heard similar stories time	16	but throughout the course of the treatment the
17	and time over.	17	weight gain meant she moved from a size 12 in
18	I hand over to Jackie to give further	18	clothes to a size 22. And the significance here
19	examples.	19	was that this further led to her isolation. She
	JACKIE: Yeah, I met one lady who was diagnosed	20	felt very embarrassed, she didn't feel she could
20		21	go out shopping. Her son said to me that she
21	ultimately with post-traumatic stress disorder	00	
21 22	as a result of the impact of her treatment,	22	felt that people judged her because of her
21 22 23	as a result of the impact of her treatment, which she described as traumatic, and that	23	weight, and her son said to me that he felt like
21 22	as a result of the impact of her treatment,		

82 1 And so she became, as I said, further 1 hepatitis C, and very few had had serious side 2 isolated, didn't go out. Again, then impacting 2 effects. 3 3 on low mood, and anxiety as well. Obviously, there was -- you know, people 4 4 As Pam said earlier, lots of families hadn't were really glad that it took less time, and 5 been informed of the likely impact of the 5 some people felt the benefits in a much quicker 6 treatment. A few people said to us they'd been 6 time frame, with one person feeling after 7 told it would be like flu-like symptoms, which, 7 a couple of weeks "I really think this has 8 as I'm sure people here know, doesn't go 8 worked", and it had. 9 9 anywhere near describing the impact of that kind With regards to people receiving this 10 of treatment. 10 treatment there was this kind of double edged 11 11 sword with this kind of "We're the lucky ones But particularly, people didn't realise that 12 the mental health and mood changes were related 12 because we're still alive and other people have 13 to the treatment, and so they went through 13 died", and actually in some situations there 14 agonies in their families in relationships, and 14 were individuals that got the feeling from some 15 self-blame and blame of each other, and some 15 of the professionals that they encountered, 16 relationship breakdowns, because they hadn't 16 "Think yourself lucky that you're getting this". 17 realised that this was not the real them, this 17 And, you know, those kind of feelings didn't 18 was what the medication was doing to them. 18 leave them in a good place, and obviously those 19 19 that hadn't had that experience were really Kay, you had some ... 20 KAY: Yeah, I was just going to talk and bring it 20 appreciative of getting a really good service, 21 forward to around 2015, and the people who have 21 and obviously clearing the hepatitis C. 22 22 been able to use the new direct acting One of the gentlemen that I met was really 23 anti-viral treatments, and many spoke positively 23 shocked that after I'd visited him early last 24 24 about this, and everyone on this treatment year, he'd had the need to go to the GP, and --25 25 programme that we've met has been cleared of was really shocked. He's had this condition 83 84 1 since the early nineties. The GP was kind of 1 together. 2 2 telling him that he'd got hepatitis C. Well, Another family I spoke to, who were hopeful 3 he's known that for a long, long time. But what 3 for a liver transplant -- it was the lady 4 was more shocking was there wasn't a discussion 4 I spoke to just recently who had the very 5 around the new treatment, and to this day this 5 significant weight gain -- her liver disease 6 man still has hepatitis C and there's 6 progressed to cirrhosis and when she was 7 a treatment available there. And that really 7 assessed for a liver transplant the assessing 8 does bring about the issue of ensuring that 8 surgeons refused to undertake the procedure 9 9 everybody is afforded the opportunity to go on because of her weight. And the family were 10 the programme in terms of the new treatment and, 10 really angry that they -- in their words, they 11 you know, that's something that was really quite 11 said she was infected by the NHS, the NHS treatment caused her weight gain, and then it 12 shocking to hear. 12 13 MS SCOTT: Jackie, some of the people you spoke to 13 was that that made her not qualify to have the discussed liver transplants with you. What were 14 14 transplant that she then needed, and she 15 you told? 15 subsequently died. MS SCOTT: Jackie, you spoke to a number of people 16 JACKIE: Yes, we saw four people who had had liver 16 17 transplants, and I wanted to just share two 17 about end-of-life care and end-of-life 18 examples of different experiences, really. 18 experiences. 19 One man had a successful transplant which JACKIE: We did. 19 20 really transformed his and his family's life, 20 MS SCOTT: Can you tell us about that. 21 and his wife said to me that it gave them nine 21 JACKIE: Again, we found a range of experiences. 22 22 of their very best years together. Sadly, the Several people sadly talked about their loved 23 23 hepatitis C couldn't be cleared and his new ones experiencing very traumatic deaths. Often 24 liver became infected, but she was very grateful 24 related to multiple organ failure. Some of them 25 for those nine years that they did have 25 were treated very appropriately and sensitively.

85 1 Others had less pleasant experiences. 1 And what she described was just that 2 One lady spoke to me about her mum not 2 contrast, within a 24-hour period, of the two 3 3 having a peaceful or pleasant passing. And different approaches, in that respect. 4 4 a few people talked to us about the cruelty, how I saw another lady whose husband had had 5 it seemed very cruel at the end of -- people 5 hepatitis C for a long time, and was -- then 6 having to have their fingers really squeezed 6 unfortunately contracted hepatitis E through 7 hard, for example, to have their blood tested, 7 food poisoning and so he became very ill very 8 et cetera. 8 9 9 One lady described to me her mum, who was They had a very poor experience in hospital, 10 dying from liver failure at the age of 51 and 10 no specialist beds were available, and he 11 11 she was in the last days of her life. And the remained in A&E. And although she was very 12 daughter was outside of the room, leaning up on 12 concerned about the decline in his condition, 13 the wall, on the wall outside of the room, and 13 she said, in an echo of the original experience 14 one of the doctors came out and he tapped her on 14 about hepatitis C, the nurses said he was just 15 the shoulder and said, "Cheer up, it might never 15 very tired. And he actually passed away during 16 happen." 16 that period. 17 And she just felt it was so cruel for anyone 17 And she tried to speak to the hospital 18 to say that, as her mum was so close to her 18 afterwards. The family complained to the 19 life. But in contrast, she then said there was 19 hospital about the quality of care that had been 20 another lovely doctor who sat with them one 20 received, and it took a year for this complaint 21 night and very gently explained that he didn't 21 to be dealt with. And the hospital said that 22 think her mum would reach the morning, and 22 they'd done all they could do and they couldn't 23 explained to them exactly what would happen over 23 find any learning from that. The family felt 24 24 the coming hours. And this felt like a really really strongly fobbed off. 25 25 And this seemed to -- this is similar to respectful and compassionate discussion. 87 88 travel on two different bus routes. It took her 1 another example that we saw, where a widow whose 1 2 2 husband had died from hepatitis C went back to certainly upwards of two, two-and-a-half hours 3 the hospital to ask if it had been caused by 3 either way. At the point that he was ending his 4 a blood transfusion which he'd had as a result 4 life, his family weren't informed that he was --5 of an injury. Obviously they found out 5 that his death was imminent. On the day that he 6 subsequently that it had. And she was asked to 6 died, his wife took her usual lengthy bus route, 7 7 sit in front of a panel of four doctors -- and and she had taken flowers to celebrate the fact 8 this is a lady who'd just been widowed, and she 8 it was their wedding anniversary, and when she 9 9 was in her late sixties/early seventies -- and arrived at the hospital, she went to the ward 10 she said -- her daughter said that they all 10 and saw the empty bed. And that's how she'd 11 backed each other up. She was there on her own 11 been told that he'd died. She's still alive, 12 against this panel of four doctors, and she just and I met with her, a very stoic lady, and a bit 12 13 felt intimidated and humiliated, really. 13 similar to what Jackie has just mentioned: hadn't got the energy and the emotional energy 14 And in both of those situations, both of 14 15 those families said the family just did not have 15 to take this matter forward. 16 the energy to carry on with those complaints, 16 So a truly tragic death for this man and his 17 because of where they were in their bereavement 17 family. 18 and coping with the condition. 18 Kay, I think you wanted to ... 19 Which leads me on to my example, which is KAY: I just wanted to move on. We've touched PAM: 19 20 not dissimilar. I mentioned earlier about the 20 on -- a number of people spoke about the impact 21 man with leukaemia who had got hepatitis B from 21 of losing loved ones to early and traumatic 22 22 deaths. And one of the families that I met had a blood transfusion, who spent the last weeks of 23 23 his life in a psychiatric ward because of the lost their child. This was a family whose 24 perceived need to nurse him in isolation. 24 teenager had had an accident, needed to have 25 His wife visited the hospital, needing to 25 a blood transfusion, and very quickly after that

90 1 transfusion she became very, very poorly, and 1 grateful that the intermediary process was 2 after a few months she died, and she was 14. 2 available for her to be able to honour her 3 3 Losing anybody is difficult. Losing a child daughter's memory and be able to have a voice in 4 is something that people will say is forever 4 this process. 5 life-changing as well, in terms of that. And 5 I think that is a really, really important 6 there was a sibling as well, and the impact on 6 part of -- you said earlier on this was a new 7 the sibling was significant. 7 process. It was so important, in particular in 8 Her voice in this process, I would like to 8 that case, and I would say in all cases, that 9 9 share with you: this option was available. 10 "There is nothing worse than the loss of 10 JACKIE: One of the things we recognised, and 11 your child. It destroys your whole life. I had 11 something that people said to us, was those 12 so much time off work. I was almost sacked. It 12 losses are tragic and traumatic for the 13 changes your personality, and it's devastating, 13 individual families, but we recognise as well 14 and you just function day-to-day, and there are 14 that those deaths were a loss to the wider 15 no words to describe it." 15 community and to society itself, really. There 16 I just want to say a little bit about this 16 were examples of the local undertaker who was 17 lady, who again, we have described about all our 17 such a support to his local community, the 18 families, was a lady -- she is now 80 years 18 much loved school lunchtime supervisor, the 19 old -- the death was quite a significant number 19 nurse who died too early at 51, all people who 20 of years ago, and she has had to 20 provided excellent services to their community 21 compartmentalise how she deals with that, to be 21 and to society overall. 22 22 able to go on for her other child and obviously One man spoke to us about the lost potential 23 23 of his brother, who died at 23. This young man the family. 24 24 But she felt so strongly for this inquiry was a talented musician and singer with 25 25 that she had a voice, and, Sir Brian, she was so excellent academic achievements. His brother 91 92 1 said: 1 therapeutic period in their lives to be able to 2 2 "I know that had he lived, he would have work through some of the issues that have been 3 made a huge contribution somehow. He studied 3 around for many, many years. 4 medicine at Oxford and then changed courses to 4 The whole issue nationally I know has been 5 study law and achieved a first class degree. He 5 addressed around mental health and stigma, and 6 was a polymath. His death at 23 meant the whole 6 I know there's lots of efforts nationally now, 7 7 country missed out on what he could have and some spoke of how positive that is, and we 8 contributed." 8 can open the doors for some of those people that 9 MS SCOTT: Kay, can I ask you then to move us on to g need that service now, even if it was 30 or 10 the next topic, which is access to psychological 10 40 years ago, as well as those more recent. 11 support. 11 Some spoke of, if it was a tailored KAY: I just wanted to start this area looking psychological support for people, it's important 12 12 13 at -- kind of historically. So looking at those 13 for people to know the history of infected blood 14 that were infected in the seventies and eighties 14 and blood products and the whole history of 15 and nineties. There was quite a common theme 15 that. Others may have commented on actually 16 with regards to there were varied examples of 16 those professionals can learn some of that away 17 those that received or were offered 17 from the therapeutic environment, but actually, 18 psychological services. And some did, but may 18 it's important that they have some knowledge 19 have had to seek that out themselves. So one of 19 of it so they can work with clients and work 20 the other things that I encountered was some 20 with them, and their experience of the 21 people I spoke to who would have found it really 21 environment. 22 22 hard to ask for help and support, and the need I'll hand over. 23 for specialist psychological services can never 23 MS SCOTT: Kay, can I just ask, were any of the 24 be underestimated, in terms of those people who 24 people that you spoke to offered psychological 25 need to have that time and that longer 25 support in the early days of the infections?

1	KAY: In the early days, for myself, that was	1	PAM: I can end this section on two positives.
2	a no.	2	The young woman I've talked about intermittently
3	MS SCOTT: No. Is that the same for the	3	with thalassaemia, she had received counselling
ļ.	JACKIE: Almost exclusively, yeah. I saw one lady	4	intermittently over the years. In the first
	who was asked to go to a group in respect of her	5	instance around the treatment around
	condition, and when she got there she was in	6	thalassaemia and how that impacted on her life,
	a group of people who had addiction problems,	7	but then, of course, as she became hepatitis C,
	and she felt that was highly inappropriate for	8	she began to have a different type of bespoke
	her, for her situation. And then another one	9	therapy around her particular needs. And that
)	where an infected person who was themselves	10	has continued, the therapy or the counsellor
1	a health professional was asked to lead a group,	11	has been flexible in the sense of being able to
2	and again, she felt that that blurred the	12	to be responsive to what she's needed at that
3	boundaries between herself as someone who was	13	time and she feels that has been a lifeline.
1	experiencing the issue.	14	My second example is quite different.
5	I did speak to one woman who had two	15	A woman who had the treatment, she'd had
3	different experiences. She was referred to an	16	interferon and ribavirin, had been through
,	NHS psychologist who told her that she must have	17	a horrendous time, one of those people who
}	had a death wish because she'd had	18	talked about it being the worst year of her life
, )	a relationship with somebody who was infected.	19	and being suicidal. It wasn't successful. She
)	But she then later sought help through the	20	then had the new treatment a number of years
1	Hepatitis C Trust, who were incredibly helpful	21	later. Her doctor, her GP, she describes as
2	to her, and supportive, and arranged for private	22	being fabulous and had talked her through how
3	counselling for her, which was excellent in	23	she needed to exercise and said, "Do 100 yards
1	taking her forward. So some very different	24	this week, do 200 yards next week, go round the
5	experiences there.	25	lake, et cetera, et cetera", to the point that
	95		96
	she felt fit and healthy, and at that point he	1	of the enhanced financial arrangements came
	and the surgery hosted a tea party for her and	2	around for the psychological impact, really
	her family to go to the surgery and have this	3	struggled in even recognising his own
	tea party. And she said again, it was		
	ted party. And she said again, it was	4	depression. His wife saw it, as did another
i	a lifeline, that the GP and the surgery got her	4 5	depression. His wife saw it, as did another family I can think of with the couple, but they
	• •		•
	a lifeline, that the GP and the surgery got her	5	family I can think of with the couple, but they
i	a lifeline, that the GP and the surgery got her through those times.	5	family I can think of with the couple, but they were struggling because they just felt so lucky
	a lifeline, that the GP and the surgery got her through those times.  So we do have these exceptional experiences	5 6 7	family I can think of with the couple, but they were struggling because they just felt so lucky they were alive that they weren't recognising
	a lifeline, that the GP and the surgery got her through those times.  So we do have these exceptional experiences from some of the people that we saw.	5 6 7 8	family I can think of with the couple, but they were struggling because they just felt so lucky they were alive that they weren't recognising their own depression and anxiety. And that had
)	a lifeline, that the GP and the surgery got her through those times.  So we do have these exceptional experiences from some of the people that we saw.  MS SCOTT: For those people that you spoke to that	5 6 7 8 9	family I can think of with the couple, but they were struggling because they just felt so lucky they were alive that they weren't recognising their own depression and anxiety. And that had manifested itself in so many ways, in particular
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lost their fathers through the infected blood

resulting in hepatitis B. And currently, there

is no financial support available for this

98 1 heard from with regards to hepatitis C had been 1 had to give up work to look after their husband, 2 told by professionals about the Skipton Fund. 2 and I think that the lady I met said, "I love my 3 3 Others found out through the press or through job but I love my husband more", and that 4 relatives, and some didn't know about it until 4 financially impacted on them but she had to give 5 it became a bigger part of the inquiry. So 5 up work because he was so poorly in terms of his 6 there was kind of a group of people, the 6 mental health and physical health. 7 majority, who did know, but there were others --7 It was common for issues such as not being 8 and I even add one gentleman who found out 8 able to get life insurance, mortgages, holiday 9 because his -- it was through reading somebody's 9 insurance. Some people didn't apply for jobs 10 story in a magazine, and then thought, "Oh, 10 because of the fear of being asked the questions 11 right, I can apply for this", and did, and 11 around their medical health and they didn't want 12 obviously was awarded. 12 to share that. 13 People spoke positively about the current 13 And I had one particular professional that 14 schemes, but all expressed anxiety that this 14 worked in the medical field that says, "I've 15 support might be reduced as part of austerity 15 been ill all my life. I'm always having tests. 16 measures, meaning that they can't rely on it for 16 My life revolved around work at a hospital as 17 long-term financial plans, and there were some 17 a professional nurse and being a patient. 18 who felt that payments were inadequate, for 18 I love my work but it affected by prospects, so 19 others, across the differing areas, different 19 I couldn't progress in my career, and it's what 20 awards, and they'd like some kind of agreement 20 I really wanted to do." 21 moving forward, that there's parity with that. 21 At the end of that section there's also, in 22 One of the things we discussed was around 22 the report, mention of the family who lost the 23 financial support, but also financial impact. 23 child not being able to afford the funeral, and 24 24 So many of the people that we spoke to had having to get a loan and borrow money to 25 25 suffered financial impact because a partner had actually be able to bury their child. 99 100 JACKIE: I met a family who had to borrow money from 1 condition. One of the dependents sadly died, 1 2 2 friends, and this was because during the period the other dependent is still alive, and very 3 that the husband was having interferon and 3 much in need of financial support, having lost 4 ribavirin treatment -- this took two years, 4 her husband, who I referred to earlier on as 5 altogether, because he had to keep having pauses 5 a self-employed businessman, and having lost all 6 to allow his immune system to recover -- and 6 access to any type of benefits and payments. 7 7 they had no income for 18 months. And they used So that's an anomaly within the system still 8 up their savings within six months. His wife 8 as well. 9 worked part-time, and that meant that they 9 MS SCOTT: The end of your report you set out the 10 weren't able to access any benefits. They 10 hopes of those you spoke to, their hopes for the 11 didn't know about the Skipton Fund or anything 11 inquiry. 12 like that, and the only help they got was Pam, I wonder whether you can just emphasise 12 13 a ten-pound reduction on their Council Tax 13 on some of those key themes emerging from that. 14 during that time. 14 PAM: Yeah, to return to where we started off, and 15 So, as I said, they had to borrow from 15 the fact that we felt very humbled and 16 families. And they found out about the Skipton 16 privileged to be part of interviewing these 17 Fund in a happenstance way as well, and 17 people and represent them at the inquiry, the 18 thankfully were able to repay their relatives 18 majority of people we know describe themselves 19 for the money that they'd loaned to them during 19 as the lucky ones because they'd survived, and 20 that period. 20 we heard this at the beginning of so many of our PAM: 21 21 Just to finish off in this section, we had interviews. 22 22 a situation for two particular families who'd Now, the interviews, the people we met, they

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were stoic, they were resilient, but also

sensitive to the situation of those who had

died, very much so, and their bereaved families

(25) Pages 97 - 100

101 102 1 and people currently who were ill and receiving 1 want the people responsible to be held to 2 treatment ... so their messages to the inquiry 2 account and to explain whether their decisions 3 3 are messages that I think the Inquiry has heard were made in good faith on the basis of what was 4 throughout its time, throughout the time you've 4 known, or whether there were other drivers such 5 been hearing evidence from those infected and 5 as finance 6 affected until this point. 6 Some were concerned about a cover up, and 7 I won't go through all of what our folks 7 wanted to know if this had happened, and who was 8 said because you've already heard them so some 8 responsible. 9 9 of the message is the same but we know as They wanted the public to know more about 10 experts by their experience, they're the 10 their experience to replace myths with facts, 11 experts, not we three, they're certainly the 11 and thereby to reduce the stigma associated with 12 experts, and they gave a very powerful 12 infections. They want the public to know that 13 collective picture revealing those common 13 they were infected through no fault of their 14 patterns and themes, and their commonality. 14 own, and that was certainly a theme throughout. 15 They'd like to see a clear timeline of when 15 Many people said that they would like the 16 infected products were first used and when they 16 financial support arrangements to be reviewed, 17 stopped, along with an understanding of how 17 to make it less of a battle for people to get 18 decisions were made along this timeline, to 18 help. They would like certainty that their 19 continue using those products, how risks were 19 current support would not be withdrawn. 20 considered, and whether this was assessed as 20 Many suggested that the attempts of 21 a risk worth taking. 21 government organisations to reach people 22 They particularly want to know what people 22 possibly infected with hepatitis C had been 23 in authority knew about the risks of infected 23 ineffective, and I think we were probably most 24 24 blood when making strategic and clinical surprised by that, that there was no reach-out 25 25 treatment decisions to continue its use. They programme for those people who had had blood 103 104 1 transfusions or received blood during those 1 a cause of death of their loved ones. 2 2 So this process, Kay already touched upon critical years. 3 They spoke about people carrying infections 3 it, has given them the strength and self-esteem 4 they do not yet know about and how the lack of 4 and the confidence, some of that confidence, 5 recall scheme following transfusions left them 5 back to be able to talk about this more openly. 6 with a ticking time bomb for years and an 6 and are very grateful for the Inquiry having 7 7 heard their voice. irreversible progression in their disease. They 8 would like to see a scheme in which everyone who 8 MS SCOTT: Is there anything else that either of you 9 received blood products or blood transfusions g two want to add before we finish? 10 from the 1970s until screening was fully 10 JACKIE: We just want to say we are so grateful to 11 effective would be offered a blood test. 11 the people who allowed us into their lives, 12 We certainly know from our work with this really, and to pay testimony to them. This has 12 13 group that the Inquiry has given them confidence 13 been a tremendous experience for us professionally and personally, and we take our 14 to speak out, and interestingly, those people 14 15 who were very, very keen to remain anonymous at 15 hats off to you folks. KAY: 16 the beginning of this process, by the end, and Yeah, I agree. 16 17 they've had contact with us, say, "I feel able 17 JACKIE: We do. 18 to now to speak out," and they feel able to now 18 SIR BRIAN LANGSTAFF: Ms Scott, do you wish to check 19 to be identified. 19 with those behind you whether anyone has any 20 I'm not suggesting that we identify those 20 further questions? 21 people today, but you can see, those of you that MS SCOTT: Yes. 21 22 22 looked at our report, some people have I think that's a no, sir. 23 identified their loved ones who have died, 23 SIR BRIAN LANGSTAFF: Well, I have one. It's been 24 because they feel able, now, to be identified as 24 said to me in a number of statements that the 25 people who have had hepatitis B, C, or HIV, as 25 author has lost trust in the doctors and

	105		106
1	clinicians who have treated him and, therefore,		ne, but it is about reaching
2	finds it difficult to accept or take medical	•	people getting access to
3	treatment now. I think one person has described		ervices because they filled
4	it as the single greatest loss arising out of	4 out the form and tic	ked the box that's in there.
5	the infected blood events.	5 I think it's that two-	way process of reaching
6	What, if any, reflection did you have of	6 out to those that we	e know about.
7	that idea from those that you were speaking to?	7 JACKIE: I saw a famil	y whose son had had leukaemia
8	KAY: I think, Sir Brian, it's around building	8 in his early childhood	od when he was six, five or
9	that trust and confidence for people to be able	9 six. And had a bloc	od transfusion as a result of
10	to step into now, in particular the gentleman	10 that which resulted	in hepatitis C infection and
11	who has still not had treatment, step into an	11 then obviously all o	f the treatment thereafter,
2	environment where some trust can be built to	12 and he wasn't even	able to see me. He had such
3	work towards eventually going and having some	13 a mistrust of anybo	dy that he saw associated
4	treatment.	14 with authority or an	ything formal. And I had to
5	I found it absolutely heartbreaking that	15 see his parents, wh	no very eloquently described
6	there's a treatment that somebody can have but	16 his situation, but the	ey said that his lack of
7	because of their experience, they won't have it.		much broader avenues of his
8	And that was further made worse by last year's	18 life, into relationship	os, personal
9	experience for the same individual. So I think	19 relationships, as we	ell as the more formal
20	it's about building trust and confidence and the	20 relationships with p	rofessionals.
21	psychological services is an opportunity to	21 PAM: Nothing to add	
22	engage, but to really engage with individuals to		FF: Can I thank you collectively,
23	say what's the way that you would want to access		for what you have done for us,
24	some services? Because for some people it would		lunch. It's been very, very
- · 25	be group. For some people it would be		ut I wanted to thank you in
	107		108
1		1 Thank you very mu	
	particular for three things. The first and	1 Thank you very mu	
2	particular for three things. The first and perhaps most important is for giving people	2 KAY: Thank you.	
2 3	particular for three things. The first and perhaps most important is for giving people a voice who would not otherwise have had one,	<ul><li>2 KAY: Thank you.</li><li>3 JACKIE: Thank you.</li></ul>	
2 3 4	particular for three things. The first and perhaps most important is for giving people a voice who would not otherwise have had one, and by goodness you've given them not only	2 KAY: Thank you. 3 JACKIE: Thank you. 4 PAM: Thank you.	ch indeed.
2 3 4 5	particular for three things. The first and perhaps most important is for giving people a voice who would not otherwise have had one, and by goodness you've given them not only a voice but a powerful one. Thank you.	<ul> <li>2 KAY: Thank you.</li> <li>3 JACKIE: Thank you.</li> <li>4 PAM: Thank you.</li> <li>5 SIR BRIAN LANGSTAR</li> </ul>	ch indeed. FF: We'll take a slightly earlier
2 3 4 5 6	particular for three things. The first and perhaps most important is for giving people a voice who would not otherwise have had one, and by goodness you've given them not only a voice but a powerful one. Thank you.  Secondly, for the empathy which you have,	<ul> <li>2 KAY: Thank you.</li> <li>3 JACKIE: Thank you.</li> <li>4 PAM: Thank you.</li> <li>5 SIR BRIAN LANGSTAR</li> <li>6 break for half an ho</li> </ul>	ch indeed. FF: We'll take a slightly earlier our or so. Shall we say
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	109		110
1	No need for me to introduce them	1	MS EDWARDS: As you know my name is Sian Edwards.
2	individually because their names are in front of	2	I am here I've been a nurse in as an HIV
3	them where they will sit. Could they please	3	specialist nurse since 1986, in various
4	come forward and be sworn.	4	countries in the world, and much of that nursing
5	PROFESSOR MYFANWY MORGAN (affirmed)	5	has also been education of healthcare workers,
6	DR NICKY THOMAS (affirmed)	6	and nurses specifically. I worked for 5 years
7	DAME LESLEY FALLOWFIELD (affirmed)	7	as an no, as a community haemophilia nurse at
8	MS SIAN EDWARDS (affirmed)	8	St Thomas', and I followed that up with two
9	PROFESSOR JOHN WEINMAN (affirmed)	9	research projects looking at life histories of
10	Examined by MS RICHARDS	10	people who had HIV and haemophilia, and also the
11	MS RICHARDS: Can I ask each of you to just briefly	11	partners, the siblings, the children, and
12	introduce yourselves and explain what your	12	parents of those who had died from HIV, who had
13	qualifications and areas of expertise are,	13	had haemophilia. I'm now a research nurse in
14	starting at the far end with Professor Weinman	14	Australia in HIV.
15	and then moving through the table.	15	DAME LESLEY: Hello, I'm a professor of psychosocial
16	PROFESSOR WEINMAN: Okay, I'm a professor of	16	oncology, which means psychology as applied to
17	psychology as applied to medicine, so I'm	17	cancer, and I am the director of a health
18	someone trained in psychology, originally in	18	outcomes research group at Brighton and Sussex
19	clinical psychology, but more recently in health	19	Medical School. This group looks at the quality
20	psychology, and so my areas of expertise are	20	of life issues for patients undergoing different
21	very much about understanding the psychological	21	cancer treatments. We also develop psychosocial
22	impact of illness and healthcare and treatment	22	support services for patients and their
23	on the individual and how people cope with and	23	families, coping with life threatening illness,
24	adjust to illness, and how that impacts on	24	and importantly and relevant to this Inquiry, we
25	recovery or outcome.	25	do a lot of work developing communication skills
1	111 training programmes for healthcare	1	112 been in terms of eliciting the patients' and
2	professionals, and as a part of that, sometime	2	families' perspectives and experiences to inform
3	ago I worked very closely with National Patient	3	the development of interventions and to assess
•	ago i worked very closely with reational i attent	0	the development of interventions and to assess
Λ	Safety Agency on designing a programme to run	1	the outcomes of interventions for nationts, and
4	Safety Agency on designing a programme to run	4 5	the outcomes of interventions for patients, and
5	for healthcare professionals on how they should	5	I've also been involved in developing methods of
5 6	for healthcare professionals on how they should communicate with patients when there was medical	5 6	I've also been involved in developing methods of research in order to better further this
5 6 7	for healthcare professionals on how they should communicate with patients when there was medical error.	5 6 7	I've also been involved in developing methods of research in order to better further this process.
5 6 7 8	for healthcare professionals on how they should communicate with patients when there was medical error.  DR THOMAS: My name is Nicky Thomas, and I'm	5 6 7 8	I've also been involved in developing methods of research in order to better further this process.  MS RICHARDS: Thank you.
5 6 7 8 9	for healthcare professionals on how they should communicate with patients when there was medical error.  DR THOMAS: My name is Nicky Thomas, and I'm a consultant health psychologist at Guy's and St	5 6 7 8 9	I've also been involved in developing methods of research in order to better further this process.  MS RICHARDS: Thank you.  Some of you I know attended hearings last
5 6 7 8 9 10	for healthcare professionals on how they should communicate with patients when there was medical error.  DR THOMAS: My name is Nicky Thomas, and I'm a consultant health psychologist at Guy's and St Thomas'. I recently retired as head of	5 6 7 8 9 10	I've also been involved in developing methods of research in order to better further this process.  MS RICHARDS: Thank you.  Some of you I know attended hearings last year where you heard firsthand some of the
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	113		114
1	quite sure how, as convener. And what we did	1	I'm going to ask you about based upon the themes
2	was to first of all have a telephone conference	2	and issues explored in your report: psychosocial
3	where we all looked at the we were asked	3	impact, social impact, stigma and
4	a whole series of questions, where we looked at	4	discrimination, loss of trust and communication,
5	the questions and thought about the expertise we	5	and care and support. And we'll broadly follow
6	had, and a sort of mixture of joint decision	6	that order.
7	making and a bit of cajoling from me led us to	7	And so I wanted to start, perhaps by asking
8	end up with taking charge of different areas of	8	Professor Weinman but, please, anyone else who
9	the report, which we've sort of indicated.	9	feels able to, please chip in at any stage with
10	MS RICHARDS: But ultimately it is a collective	10	any of the questions, to ask you about what
11	report?	11	you've set out in section 13.1 of your report
12	PROFESSOR WEINMAN: Very much so, yeah.	12	under the heading "Psychological impacts of
13	MS RICHARDS: You've all looked at all of it?	13	infection on people infected and affected".
14	PROFESSOR WEINMAN: Yeah.	14	You've said, Professor Weinman, in that
15	MS RICHARDS: And as I understand it there are no	15	section, that:
16	areas of disagreement professionally between	16	"All long-term illnesses have psychological
17	you?	17	impacts on those with the illness, and any one
18	PROFESSOR WEINMAN: No.	18	illness will have effects on an individual's
19	MS RICHARDS: You're also going to be producing	19	emotional state and their self-identity, as well
20	a supplemental report responding to what's	20	as giving rise to adjustment to the pain,
	described as part 2 of the questions in	21	
21	•		incapacity and demands of long-term treatment
22	a supplemental letter of instruction dated	22	and lifestyle change."
23	23rd December.	23	I wondered if you could elaborate on that
24	PROFESSOR WEINMAN: Correct, yes.	24	for us please.
25	MS RICHARDS: Now there are essentially five areas	25	PROFESSOR WEINMAN: Well, for any of us, an illness
	115		116
1	or, indeed, any threat to our health is a sort	1	116 depression, really serious psychological impact.
1		1 2	
	or, indeed, any threat to our health is a sort		depression, really serious psychological impact.
2	or, indeed, any threat to our health is a sort of threat to one's sense of self and who you	2	depression, really serious psychological impact. But all of that really will be very much
2	or, indeed, any threat to our health is a sort of threat to one's sense of self and who you are, and your place in the world. And anybody	2 3	depression, really serious psychological impact. But all of that really will be very much governed by how much the illness impacts on
2 3 4 5	or, indeed, any threat to our health is a sort of threat to one's sense of self and who you are, and your place in the world. And anybody who, you know, either has sudden onset or slow onset or whatever of a condition, tries to make	2 3 4	depression, really serious psychological impact. But all of that really will be very much governed by how much the illness impacts on one's sense of self and one's self-identity. It's absolutely crucial. For some people they
2 3 4	or, indeed, any threat to our health is a sort of threat to one's sense of self and who you are, and your place in the world. And anybody who, you know, either has sudden onset or slow onset or whatever of a condition, tries to make sense of it. In the first place they need to	2 3 4 5	depression, really serious psychological impact. But all of that really will be very much governed by how much the illness impacts on one's sense of self and one's self-identity.
2 3 4 5 6 7	or, indeed, any threat to our health is a sort of threat to one's sense of self and who you are, and your place in the world. And anybody who, you know, either has sudden onset or slow onset or whatever of a condition, tries to make sense of it. In the first place they need to try to understand what's going on.	2 3 4 5 6	depression, really serious psychological impact. But all of that really will be very much governed by how much the illness impacts on one's sense of self and one's self-identity. It's absolutely crucial. For some people they are able to somehow negotiate that and work around the illness and accommodate the illness.
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117 118 1 treatment, you know, reaching out to people for MS RICHARDS: And that which you've described, that 2 support and so on. There are a whole -- very 2 all long-term illnesses have psychological 3 3 complex number of processes, and that happens to impacts in the way that you've touched on, that 4 everybody, regardless, and I think obviously in 4 is not as I understand it any kind of a new 5 the case of people who already had an illness 5 idea; that's something that's been known and 6 and then were then infected further with, you 6 understood for many years. 7 know, a blood product, then the process becomes 7 PROFESSOR WEINMAN: Yeah, very much so. I think 8 even more complicated. 8 what's new is that we understand more about some 9 9 Now you're dealing not just with one, of those ways that people think and ways that 10 sometimes two and sometimes three conditions, 10 people cope. And particularly ways that they 11 with a whole range of treatments. And the 11 can be supported through an understanding of 12 having to adjust to things like pain, incapacity 12 those very individual processes. Yeah. 13 and treatments which sometimes can be truly 13 MS RICHARDS: Can I ask you a little more about what 14 horrible, as we've heard many times in these 14 it is about having long-term illnesses that can 15 hearings, takes a huge toll on the individual 15 have the consequences you describe. You've touched on some of them already. 16 and those around them. So I think that's -- you 16 17 know, what you've really summarised from that, 17 First of all, pain. The psychological 18 you know, the review on how people cope with 18 impact of coping with pain. What can you tell 19 illness, involves all those processes which are 19 us about that? 20 complex, they're not fixed in time, they change 20 PROFESSOR WEINMAN: There is a huge amount that's 21 over time, as new events emerge, as people 21 known about, you know, the impact of pain on 22 perhaps who you were relying on become less 22 individuals. We know that -- because, in a way, 23 reliable and other people become important in 23 pain is a very hidden thing, you know, that 24 your life. So the whole social impact as well 24 people who are experiencing it experience it 25 25 was very much part of that process. hugely -- but pain is not out there, it's not 119 120 1 like a broken arm or whatever, that often 1 incapacity. So the individual who is often 2 I think we know that people are having to cope 2 suddenly, but sometimes gradually, no longer 3 with pain very much on their own, with other 3 able to do that which they could previously do. 4 people perhaps not realising the extent of what 4 How does that bear upon the psychological 5 they're having to cope with. We know that 5 experience of the individual? 6 people cope with pain in many, many different 6 PROFESSOR WEINMAN: It's absolutely central, in the 7 7 ways. There are questionnaires and assessments sense that -- I mean, disability sort of works 8 which look very specifically at pain coping and 8 at two levels, in a way: what the individual can 9 9 particularly how pain in itself activates a lot and can't do, and then how that can impact on, 10 of those processes I was talking about earlier, 10 again, their identity or as a competent person 11 the different sorts of thoughts and fears and 11 or as someone who can do things, work, be 12 worries, about oneself and one's future. And a parent, whatever it is, all those roles, 12 13 pain can generate a huge amount of fear. You 13 disability impacts all those roles. But it's 14 know, fear of the future, fear of what you can 14 also very much something -- and again we'll hear 15 and can't do, and so on. 15 about this more I think particularly from 16 So pain itself, although one can think of it 16 Myfanwy -- something which occurs in a social 17 in quite biomedical terms, as an experience it's 17 setting. People are often defined by their 18 a very individual and very demanding process, 18 disability because of things they can't do, out 19 which takes -- people cope with in many, many 19 there in the real world, and the limitations 20 different ways. And I know that when -- later 20 that are placed upon them. 21 on we hear about support for patients, you know, 21 So again, it's, you know, extremely 22 22 it's very much -- something I think Nicky will pervasive. It's something which has a very 23 23 strong impact, both on people's moods, their talk about. 24 MS RICHARDS: One of the other factors you touched 24 emotions, so it's not surprising that people 25 on, both in your report and in your answer, is 25 with disabling conditions are often very

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1	psychologically affected by those.	1	area which John has identified as around the
2	But also, you know, it's very much affected	2	notion of being a competent person. So you may
3	by the attitudes of other people and the support	3	have this chronic illness but you are still
4	they get from other people.	4	a mother, a father. You're also maybe holding
5	I don't know whether, Sian, you wanted to	5	down a job and you are still expected to hold it
6	sort of add to that, really?	6	down. So you're managing multiple roles. And
7	MS EDWARDS: Well, I think whenever we talk about	7	how do you move from those roles in a seamless
8	things like this, that we talk about individuals	8	fashion and then manage a chronic illness as
9	and different people react in different ways to	9	well? Something which is very disabling, like
10	these issues, but the support they get, as we've	10	pain and distress. So I think that's a really,
11	heard, from health services, from family, from	11	really big thing, of helping people to move
12	friends, from social groups like churches, are	12	through, and if the support isn't there, then it
13	all fundamental on how they're going to be able	13	becomes really overwhelming, and difficult.
14	to cope, and where those groups or those	14	MS RICHARDS: And how does incapacity potentially
15	services are not there for them, then you get	15	bear on the concept that you touched on, the
16	this idea of separation and isolation and trying	16	concept of self-identity?
17	to cope with this pain, this incapacity alone.	17	PROFESSOR WEINMAN: Massively, massively. Because
18	And we know that dealing with things alone has	18	self-identity is quite a complex things, but
19	very major psychological impacts on people, on	19	essentially what it boils down to is how you
20	a long-term basis. And of course, when many of	20	think about yourself, your sense of self-worth,
21	the things that we have been hearing and talking	21	and as Nicky just said, that can be based around
22	about, many of these people were children,	22	different you know, we all have different
23	adolescents and young people, as well as adults.	23	selves, you know, parents, workers, brothers,
24	DR THOMAS: If I can add to that, in terms of my	24	sisters, whatever, and incapacity can erode any
25	work as a clinician, I think a really important	25	of those bits of oneself in all sorts of very
1	123 striking ways, and I think we heard very much	1	One of the sort of crucial components of
1 2	striking ways, and I think we heard very much	1 2	One of the sort of crucial components of
2	striking ways, and I think we heard very much from well, both our intermediaries today and	2	One of the sort of crucial components of that is something we call timeline, so your
2	striking ways, and I think we heard very much from well, both our intermediaries today and also from what we read in terms of people just	2 3	One of the sort of crucial components of that is something we call timeline, so your sense of how long is this going to last. If you
2 3 4	striking ways, and I think we heard very much from well, both our intermediaries today and also from what we read in terms of people just feeling they were no longer, you know, that	2 3 4	One of the sort of crucial components of that is something we call timeline, so your sense of how long is this going to last. If you have an uncertain condition, one that flares,
2 3 4 5	striking ways, and I think we heard very much from well, both our intermediaries today and also from what we read in terms of people just feeling they were no longer, you know, that person, that parent, you know, that worker. So	2 3 4 5	One of the sort of crucial components of that is something we call timeline, so your sense of how long is this going to last. If you have an uncertain condition, one that flares, one that maybe responds to treatment but where
2 3 4 5 6	striking ways, and I think we heard very much from well, both our intermediaries today and also from what we read in terms of people just feeling they were no longer, you know, that person, that parent, you know, that worker. So that erosion can take place.	2 3 4 5 6	One of the sort of crucial components of that is something we call timeline, so your sense of how long is this going to last. If you have an uncertain condition, one that flares, one that maybe responds to treatment but where maybe treatment makes things worse, that whole
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	125		126
1	told that there was no treatment available,	1	those is Professor Deborah Christie who works
2	being told that they were likely to die within	2	with children and adolescents at Great Ormond
3	a very short period of time. How does that	3	Street and I think in the report some of the
4	impact both upon the sense of self-identity and	4	parts she's written about are very much either
5	the broader psychological condition of the	5	about, you know, parents having to protect
6	individual?	6	children from that dreadful, you know,
7	PROFESSOR WEINMAN: I mean one of the reasons why	7	prognostic information, or when the children or
8	people develop that way, their own way of	8	adolescents had that, and I certainly heard this
9	thinking about an illness is to develop some	9	in one of the hearings I attended, when you have
10	sense of mastery and control, to get that back,	10	that in your heads, that the sense of well,
11	because that what goes, you know, when you	11	giving up, not making an effort, that just, you
12	receive that sort of information.	12	know, the lack of motivation to do anything
13	And obviously being given that sort of, you	13	becomes huge. Because you end up, you're very
14	know, very clear, you know, that you haven't got	14	likely to end up thinking, "Well, whatever I do,
15	long to live or, you know, you have	15	I have no control. This is going to happen.
16	a life-threatening illness, absolutely really	16	This is inevitable", so that sort of fatalistic
7	impacts on that control, which is very	17	view, if you like, again is very undermining in
8	important. Control is really important in all	18	terms of one's motivation, one's sense of
9	our lives, control is important whatever we're	19	future, one's sense of self.
20	doing. But in the context of illness that	20	So again, very big effects.
21	really eats into that process hugely.	21	MS RICHARDS: Dr Thomas, I see you nodding.
22	I think the other rather crucial bit of that		DR THOMAS: I was just agreeing with John, really,
23	is the life stage of the individual, and we	23	just the control is so important, but the
24	don't we've two colleagues missing from our	24	ability to plan one's life. We all take it for
25	group today who will be here tomorrow. One of	25	granted, you know, we plan our holiday for next
	407		400
1	127 year or, you know, sort of for the summer, but	1	And actually as time goes on, that six months
2	if you have a physical disabling illness, that	2	comes and that year comes, and having to live
3	you can put the plans in place, but the rug is	3	through that channel of uncertainty is
4	going to be moved from under your feet when the	4	a constant for many people, particularly in the
5	holiday arrives and so on. Time and time again	5	early days when there was no treatment.
6	that's been experienced and it's come through in		MS RICHARDS: You've noted in the report that the
7	some of what we've heard in hearings and also in	7	cumulative effect of multiple medical problems
8	the transcripts that people's inability to plan	8	and conditions has been less well studied and is
9	just takes away the sense of, you know, living	9	less well understood.
0	life, really. You can't plan for tomorrow. You		PROFESSOR WEINMAN: Mm.
11	can't plan for next year. It's just very, very		MS RICHARDS: But that research has shown a close
2	difficult for people, I think.	12	response relationship between the number of
3	MS EDWARDS: One of the things that does come across	13	health problems a person experiences and the
	quite clearly is that lots of people were given	14	likelihood of depressive symptoms and impaired
14 15	that information when they were not actually	15	quality of life. Can I just invite you to
	ill, and I think that was key, because the		
6  7	·	16	expand on that?
17	diagnosis of being told "You're HIV positive",		PROFESSOR WEINMAN: Sure. It's quite curious, you
8	many people were given the misinformation "You	18	know, particularly now, as people get older and
	have got" for some people were told "six	19	develop many, you know, conditions, not just
	months to live" when actually at that moment	20	one, that a lot of the research in my field has
20	they were healthy individuals dealing with maybe	21	just focused on, you know, people with diabetes,
20 21			people with rheumatoid arthritis, haemophilia,
20 21 22	their haemophilia, but were given, yeah,	22	·
20 21 22 23	their haemophilia, but were given, yeah, 18 months to live, six months to live. And that	23	whatever, and that lack of research on people
19 20 21 22 23 24 25	their haemophilia, but were given, yeah,		·

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1	last year that really pointed out. And it also	1 MS RICHARDS: And you've identified in your report,
2	partly reflects the way hospitals work, people	2 as you say, in particular, four factors that
3	are treated in one department, in another	3 might bear upon the nature or extent of adverse
4	department as if they're somehow split up, and	4 psychological impact: one was severity in terms
5	it's the same person.	5 of clinical conditions and symptoms; patients'
6	Where people have looked at people with	6 perception of their condition; patients' coping
7	multiple conditions, certainly the probability	7 responses; and the quality of their support.
8	and the severity of psychological impact is	8 PROFESSOR WEINMAN: Yeah, exactly.
9	greater, not surprisingly, as people go have,	9 MS RICHARDS: We'll come on to talk about support in
10	say one or two or three conditions. So there	10 particular later.
11	is but it's not a linear effect. It's not	11 DAME LESLEY: May I just add something to that, to
12	a stepwise effect. And in fact and you might	12 pick up on what John has just been talking
13	go on to ask me this now what we do know is	about, people being treated in silos depending
14	that the actual severity of someone's condition	on one condition rather than the multiple
15	doesn't necessarily predict their psychological	15 co-morbidities that they might have, it leads to
16	response. There are a whole lot of intervening	16 another problem for them, which is that they
17	factors, many of which I spoke about at the	17 will bump into a whole plethora of different
18	beginning. You know, the way people make sense	18 healthcare professionals who will give them all
19	of it, the support they have, how they cope and	19 sorts of different and sometimes competing
20	so on.	20 information, and one of the anxieties that many
21	So broadly, you know, if we look across at	21 people find is exacerbated by this is the fact
22	a sort of population level, certainly we know	that they no longer have any continuity of care
23	people with more and more conditions are more	23 or information. And therefore, an expert in,
24	and more psychologically affected. We know	24 you know, HIV, AIDS, might give totally
25	that. But the relationship is quite complex.	25 different information to the haematologist
	131	13
1	dealing with the, you know, consequences of	13 1 is looking at there, if we go back to our sense
	dealing with the, you know, consequences of	1 is looking at there, if we go back to our sense
2 3	dealing with the, you know, consequences of haemophilia. And I think the disconnect between	1 is looking at there, if we go back to our sense 2 of one's self and the identity, people whose
2 3 4	dealing with the, you know, consequences of haemophilia. And I think the disconnect between all the different sources of information the	1 is looking at there, if we go back to our sense 2 of one's self and the identity, people whose 3 self-system has already been quite severely
2 3 4 5	dealing with the, you know, consequences of haemophilia. And I think the disconnect between all the different sources of information the patients and families get can lead to further anxiety. How can I plan, what can I do, when	is looking at there, if we go back to our sense of one's self and the identity, people whose self-system has already been quite severely challenged by having an illness, having to accommodate to an illness, and then another one
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1	bipolar affective disorder, anxiety and	1 condition, and I think, you know, in that sense
2	depression, in particular anxiety and	the whole of the report needs to be sort of
3	depression.	3 perhaps thought of here. What you're dealing
4	I'm not going to ask you to comment	4 with are people who experience massive stigma,
5	obviously on the circumstances of any individual	5 people who have where communication has been
6	or the causal link between infection and	6 poor, where they've perhaps not felt supported
7	psychiatric disorder in any individual, but is	7 or even felt rejected by the healthcare system.
8	this fair: that it's entirely plausible to think	8 I think it's the whole plethora of changes that
9	that infection with HIV, infection with	9 are really responsible for, you know, the
10	hepatitis C and the kind of circumstances that	10 likelihood that much greater likelihood of
11	we've seen described in the statements might	11 people developing, you know, serious
12	lead to the development of a psychiatric	12 psychological problems and reactions.
13	disorder of that kind?	13 MS EDWARDS: And then add treatment.
14	PROFESSOR WEINMAN: Yeah, I mean we know that for	14 PROFESSOR WEINMAN: And treatment, yes.
15	any major long-term health problem, a proportion	15 MS EDWARDS: You add the treatment and specifically
16	of people will develop, you know, major mood	16 these treatments, particularly interferon and
17	change. Typically anxiety in the early stages	17 ribavirin, which are debilitatingly the
18	but often leading to depression. So, you know,	depression is of an extreme type, so as John
19	a group of patients I know well, patients with	19 said you're dealing with stigma, you're dealing
20	rheumatoid arthritis, something like 20, 25% of	20 with loneliness, you're dealing with isolation.
21	those patients we know will develop those major	21 So the amount of factors that actually result in
22	mood problems, because of the changes that are	22 a depressive and sometimes serious psychotic
23	brought about by the condition.	23 illness is not surprising.
24	But I think what we're dealing with here is	24 <b>DR THOMAS</b> : Yeah, and just to add that the
25	not just the psychological reaction to the	25 traumatising experience of multiple contacts
1	135 with a very distrustful healthcare system, where	136 1 well as in obviously the group of people we're
2	you go and, as we've heard, going to many	2 talking about here.
3	different departments for treating of various	3 I don't know, I mean you're much more
4	problems and that in itself is very, very	4 experienced with that group of patients's.
5	traumatic. So you can see how, again, it adds	5 MS EDWARDS: Yeah, HIV is a debilitating illness
6	to the complexity and makes the depression move	6 with some very difficult treatments to take.
7	into another area.	7 But added on is that stigma. You add on to that
		_
8 0	MS RICHARDS: Other psychological or emotional	8 the haemophilia, you add on the hep C, you add
9 10	impacts described by infected individuals to the	9 on the hep C treatments, and I think as John
10	inquiry, again in written and oral evidence,	alluded to earlier, sometimes it's not the
11	include mood swings, panic attacks, fear, anger,	severity of the illness that causes the problem,
12	paranoia, loss of confidence, tearfulness,	it's the perception of the illness and the
13	insomnia, nightmares, loss of libido, anhedonia,	13 self-perception. And many people that we read
14	hopelessness, despair, shock.	and in my own experience, it was the association
15	Those are all, again the kind of responses	with as we heard earlier on, I needed to say who
16	that you would understand and expect, given the	16 I was and how I became infected, and we live in
17	range of different factors and circumstances	17 a world, and certainly we did live in a world
	you've identified.	that was more severely stigmatising than we do
	PROFESSOR WEINMAN: Very much so and I think, you	even today, with regard to homosexuality and
19		20 also towards drug users, and when these are
19 20	know, we know quite a lot about the	
19 20 21	psychological responses of people with, for	21 associated with these illnesses you're dealing
19 20 21 22	psychological responses of people with, for example, we know quite a lot about psychological	22 with an awful lot of factors.
19 20 21 22 23	psychological responses of people with, for example, we know quite a lot about psychological responses of people with HIV and AIDS, that's	<ul><li>with an awful lot of factors.</li><li>MS RICHARDS: Can I ask about the psychological</li></ul>
18 19 20 21 22 23 24 25	psychological responses of people with, for example, we know quite a lot about psychological	22 with an awful lot of factors.

	137		138
1	we've heard from multiple witnesses, of the very	1	from many of our witnesses, particularly in the
2	severe physical and mental effects of in	2	early days when some of those very harsh
3	particular the early treatments and when	3	treatments that didn't really seem to actually
4	I say "early", this was over a number of	4	even be working, and they were relentless and
5	years for HIV and for hepatitis. So in	5	went on and where symptoms didn't clear and
6	addition to coping with a life-threatening,	6	so on, and sort of, you know, people again weigh
7	serious illness, individuals then had to endure	7	up in their own minds: is it really at the
8	treatment which in many cases brought symptoms	8	end day, is it really worth me going on with
9	worse than the illness itself.	9	this?
0	How may that affect the individual's	10	So that's one set of issues. The treatments
11	psychological experience and condition?	11	themselves cause a lot of symptoms that people
12	PROFESSOR WEINMAN: One of the areas I work in I do	12	then have to cope with in a whole range of ways.
13	a lot of work in now because I'm linked to	13	And there's this terrible conflict which can
4	not only to a school of medicine, to the school	14	happen inside people's heads about, well,
5	of pharmacy, is looking at how people cope with	15	needing and wanting to take the treatment, with
6	and deal with treatments, and what we know is	16	the possibility of getting better, but then
7	that amongst people with long-term any	17	actually all the evidence that comes back, the
8	long-term major condition, around 40, 50% of	18	feedback, is that (a) either the condition
9	people actually don't decide, make a decision	19	the hep C has not cleared, or that there's no
20	not to follow that treatment long-term. That's	20	difference in the viral load, or whatever the
21	what we call non-adherence. Because in many	21	issue is and I think that set of conflicts
22	cases that treatment you know, they feel	22	that people are faced with is huge, and, you
23	worse when they're on the treatment. So it's	23	know and when we look at non-adherence to
24	sort of a decision that people can and do make,	24	treatment, which is often criticised, you know,
25	quite commonly. And obviously, what we heard	25	by a lot of my medical colleagues I work
1	139 with, and I work across a range of conditions,	1	140 subcategories of that in your report. People
2	are very critical of people who don't adhere to	2	completing the treatment, enduring it for
3	treatment, "Here we're offering them the	3	prolonged periods of time, only for the
4	possibility of getting better, and they're not."	4	treatment to be unsuccessful. And what's that
5 6	And we know, you know, from huge reports,	5	experience, potentially, like for an individual
	WHO and otherwise, that many make that choice	6 7 5	psychologically and emotionally?
7	because they don't in the end either don't		ROFESSOR WEINMAN: I mean, truly awful, because yo
8	believe in the treatment also issues about	8	have you've struggled with that terrible
9	not trusting the provider of that treatment,	9	conflict and you've stuck with it and stuck
10	which I know we'll get on to and so people	10	with it in the belief and, you know, beliefs
11	are then left with, you know, sometimes a bit of	11	are huge in illness, beliefs in treatment,
	a sense of guilt, "I'm not really doing what	12	beliefs about the nature of the condition and so
13	I should be doing", sometimes the people around	13	on in the belief that ultimately you're going
13 14	I should be doing", sometimes the people around them can be critical because they may not be	13 14	to get the benefit. So you've struggled for
13 14 15	I should be doing", sometimes the people around them can be critical because they may not be thinking that the person is making sufficient	13 14 15	to get the benefit. So you've struggled for a long period of time. So when it turns out
3  4  5  6	I should be doing", sometimes the people around them can be critical because they may not be thinking that the person is making sufficient effort with the treatment and so on.	13 14 15 16	to get the benefit. So you've struggled for a long period of time. So when it turns out that actually, nothing has changed or sometimes
3  4  5  6	I should be doing", sometimes the people around them can be critical because they may not be thinking that the person is making sufficient effort with the treatment and so on.  So it's quite a complex set of issues,	13 14 15 16 17	to get the benefit. So you've struggled for a long period of time. So when it turns out that actually, nothing has changed or sometimes things have actually become worse, then that is
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12 113 14 115 116 117 118 119 220 221 222 223 224	I should be doing", sometimes the people around them can be critical because they may not be thinking that the person is making sufficient effort with the treatment and so on.  So it's quite a complex set of issues, there, apart from the actual psychological impact, as you say, the pain, the side effects and other aspects of treatment, there are a whole load of issues about having to continue with something which doesn't seem to be doing	13 14 15 16 17 18 19 20 21 22	to get the benefit. So you've struggled for a long period of time. So when it turns out that actually, nothing has changed or sometimes things have actually become worse, then that is a really devastating, having gone through that struggle.  And I mean, you know, even when I don't know, maybe you're going to come on to this, but even when people have been through a major

1	141 working in cancer that people can still be	1	people, even if they do get some respite from
2	left with quite major doubts about whether this	2	the disease, isn't showing signs of progression,
3 4	thing has really gone away. You know, because their experience is still there. They might be	3 4	nevertheless can't enjoy that sort of freedom or the fact that some of the tests look quite
			•
5 6	getting symptoms and so on. So and I think we referred to that as a fear of recurrence.	5	positive, because they are so anxious about when it's going to actually re-emerge. And that
7	People worrying that somehow you know, we're	7	becomes paralysing in terms of enjoying life and
8	not experts on our own bodies, we don't know	8	what is in front of you.
9	what's happening inside us, so this whole notion	9	MS EDWARDS: The other thing about particularly
9 10	that there could still be something in there,	10	these treatments with interferon and ribavirin
11	that there could still be something in there, that fear of recurrence, which is quite	11	is that for many people, they didn't have
12	a crippling, quite a ruminative sort of process,	12	symptoms, and so their illness could be it
13	can really stop people in their tracks as well.	13	could remain anonymous. You didn't have to tell
14		14	
	DAME LESLEY: It also makes people hyper vigilant		anybody. But once you start a treatment like
15 16	about any symptoms as well. I think one of the there's two other difficult things here,	15 16	that, you are so unwell, and often people weren't aware they were going to be that sick.
10 17	I think, that we're we ought to touch on,	17	And it changes everything: your work, your
	which is that, certainly at the beginning of	18	parenting, everything in your life, if you are
18 19	many of the treatments, the likely benefits were	19	that unwell with the fatigue and depression that
20	very uncertain. Medicine anyway is not the	20	comes. That means that you then have to
21	certain science we'd love it all to be. And	20	disclose the reason for your ill health and so
22	those treated sort of in the early stages, when	22	there's major complexities to some of these
23	people were offering them some, they had very	23	treatments and the psychological impact that
23 24	little evidence that it would be beneficial.	23	they have, because you can't hide it any more.
2 <del>4</del> 25	And as John says, it's the same in cancer. Some	25	MS RICHARDS: We'll come back to the impact of loss
1	143 of trust and poor communication and stigma and	1	support but they were very much in a minority,
2	discrimination, but I just wanted to ask you	2	and, I think so then school becomes a really
3	about one observation you made in your report on	3	aversive experience for individuals. Not only
4	page 2. You've talked in this context of the	4	is it hard going, you have patchy attendance so
5	negative effects on academic behavioural and	5	you fall behind. So there are a whole lot of
6	social adaptation, and I wondered if you could	6	social effects, if you like, of the, you know,
7	elaborate upon that, please.	7	the symptoms that get in the way of, you know,
8	PROFESSOR WEINMAN: I think there's sort of two or	8	consistent schooling and so people are falling
9	three main elements to that, really. One is	9	behind with all the negative reactions that
10	that and obviously this is we're looking at	10	follow.
11	school-aged individuals and when you, you	11	I think that's one aspect. But the other is
12	know, what we heard from, certainly from parents	12	that for some of those treatments, and
13	and from, you know, people who have been	13	certainly well, we know with HIV, HIV can
14	infected, that when you feel so bad, you know,	14	have neuro-psychological effects. It can impact
15	you're fatigued, you've got pain, you've got	15	on brain function in a range of ways, but
16	symptoms and so on and you're feeling nauseous,	16	certainly a lot of those very harsh early
17	(a) you don't want to be at school so we know	17	treatments also had, you know, negative effects
18	there are a lot of school absences and, again,	18	on cognitive functioning, the ability to
19	we see this in other conditions, when you're at	19	concentrate, to remember, to problem solve. All
19 20	school, you're not you can't really engage	20	those basic skills that one needs to assimilate
			school work from and so on.
21	that well. Sadly, children pick that up. Those	21	
. ). ,	people often are not supported unless, you know,	22	So I think, you know, at a functional level
	fellow children are well informed.	23	and at a social level, school becomes very
23	I magnithere are some good examples of where	24	challonging for a lot of individuals. I think
22 23 24 25	I mean there are some good examples of where schools did make a huge effort and provided	24 25	challenging for a lot of individuals. I think those they're not separate because those

4	145		146
1	things interact very much, but both have		PROFESSOR WEINMAN: It's something that's, you know
2	profound effects on people's ability to both	2	widely reported in a lot of different areas, not
3	engage and benefit, and process all the	3	only in illness but also in military situations
4	information that one needs, because, you know,	4	and so on, that people and I think we heard
5	whatever we think about schooling, it is very	5	it a little bit from our intermediary panel
6	much a time when we have to process a lot of	6	earlier that people who are in a dreadful
7	information and do stuff and keep up.	7	situation but survive it, feel very often
8	I think that becomes very hard when all	8	feel very bad about those, you know, who didn't
9	those other negative things are going on.	9	make it. And there's that whole whereas
10	MS RICHARDS: You've picked up in this section of	10	often, you know, when something awful happens to
11	your report on evidence that you read or saw	11	us we go through the "Why me? Why has it
12	about the effects of watching others die, and we	12	happened to me", and that whole self blame. But
13	know from the evidence that we've heard, that in	13	here the self-blame is of a very curious sort.
14	the absence of formal support networks, people	14	It's almost blaming yourself for surviving and
15	formed their own support networks through	15	feeling bad that you weren't part of the group
16	a range of different groups or organisations.	16	that were left behind.
17	People within the haemophilia community knew	17	I mean it's a very powerful set of emotions
18	each other through attending centres or through	18	that can be very debilitating. But because it's
19	attending Treloar School and you've picked up	19	sort of blame, a sense of blame towards oneself
20	and quoted in your report the experiences of	20	it's a very potent source of depression, and we
21	those going to multiple funerals. I wanted to	21	know that when people experience that, that the
22	ask you two aspects, about two aspects of the	22	likelihood of becoming depressed is very high.
23	psychological implications of that.	23	I don't know, Sian, if you because you
24	The first is survivors guilt. Could you	24	must have seen a lot of people in that
25	explain that concept further for us.	25	situation.
4	MC EDWARDS. Fecentially you so to a support group	4	148
1	MS EDWARDS: Essentially you go to a support group	1 2	self-identity again.
1 2 3	<b>MS EDWARDS:</b> Essentially you go to a support group for support, and friendship, and to share your		self-identity again. <b>PROFESSOR WEINMAN</b> : Yeah, massive increase in
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	149		150
1	looking from a psychosocial perspective at the	1 DAME LESLEY: Yes (overspeaking)	
2	experiences of those infected through blood and	2 <b>DR THOMAS</b> : I spoke to a psychologist in the field,	
3	blood products, and you've drawn attention	and they also didn't have anything to recommend	
4	I won't take time with putting it on screen, but	4 as well. So it's very poor, poorly researched.	
5	it's now part of the Inquiry's material to	5 MS RICHARDS: One particular feature from the	
6	one short research paper by Riva and others from	6 evidence that the Inquiry has heard, and this	
7	2017. Are you aware of whether there's been any	7 may be particularly relevant to the concept of	
8	broader research, whether from a psychological,	8 self-identity that you have described, is that	
9	psychosocial or psychiatric perspective into	9 for many people their whole lives have become	
10	these issues, specifically in relation to those	10 defined by the condition with which they were	
11	infected through blood or blood products?	11 infected, and treatment for it, the symptoms,	
12	PROFESSOR WEINMAN: I tried very hard to find a good	12 and so on. But the result that they have had to	
13	research base for that and it wasn't there.	13 live a life completely different from the life	
14	I contacted journal editors. For example I know	14 that they would otherwise have expected to do,	
15	the editor of a big American journal called	15 and they have lost opportunities, been unable to	
16	HIV Behaviour, and asked him to go back through	16 fulfil a potential that would otherwise have	
17	his records. I looked very widely that Riva	17 been there. Again, how does that bear upon	
18	paper, it's not really a paper, I mean, it's	18 their psychological experiences?	
19	a sort of an overview on some of the	19 PROFESSOR WEINMAN: Again, hugely, because the	ne
20	psychological consequences, the	20 going back to what I was talking about earlier,	
21	neuropsychological effects and then looking at	21 about one's sense of self, one's sense of	
22	medical and legal issues, so it's not even an	22 self-identity, I think some people have said,	
23	empirical paper. So I don't know whether	23 you know, if you think of illness as a sort of	
24	colleagues had similar we've had very hard	24 set of things and oneself as a set of things,	
25	you know it's	and for some people, you know, the sense of	
	151		152
1	151 oneself can be completely obliterated and	DAME LESLEY: But there are internal processes the	
1 2	oneself can be completely obliterated and I think we heard this by you know, because	DAME LESLEY: But there are internal processes the     give you yourself identity but I think one of	
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2	oneself can be completely obliterated and I think we heard this by you know, because day-to-day, this is what's happening to you, there is nothing else, so those multiple selves,	give you yourself identity but I think one of the things that I think we were all struck by were the external processes that defined	at
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2 3 4	oneself can be completely obliterated and I think we heard this by you know, because day-to-day, this is what's happening to you, there is nothing else, so those multiple selves,	give you yourself identity but I think one of the things that I think we were all struck by were the external processes that defined somebody as being "the AIDS boy", you know, "the woman with hep C", and sometimes they assume	e e
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1	have better support and so on. So there is huge	1	survive and who made their life decisions
2	variation, I think we need to stress that	2	accordingly, but who find themselves several
3	variation. And I think we also alluded to the	3	decades on still here.
4	fact and I've seen this in quite a number of	4	What could you tell us about the experience
5	major illness groups some people go on	5	of people who might fall within that category?
6	actually to derive benefit from having gone	6 N	IS EDWARDS: I think throughout the HIV patient
7	through that process, and that process of	7	group specifically, there was so much
8	so-called benefit finding, seeing, you know, the	8	information in the early days about death,
9	sort of silver lining, if you like, in this	9	"You're going to die. Spend your money. Don't
0	horrible cloud, is something that can really get	10	bother going to school, you know, you have two
1	people through. And it's not that uncommon, you	11	or three years to live. Live it. Live it well,
2	know, certainly in the cancer literature, in the	12	because you're not going to be here".
13	heart disease literature, is benefit finding.	13	Many of those people did die, but there are
4	You know, you go through something dreadful but	14	many people, who are sitting here today, who,
5	you emerge from it stronger, you do other	15	you know, come the late 1990s, and treatment was
6	things, you cope well.	16	at last effective, who although that was a huge
7	It really goes back to this huge variation,	17	celebration in many ways, it was a major
8	and that we must not generalise. Yeah.	18	readjustment, major.
9	MS RICHARDS: Can I ask, you have talked about the	19	People had not planned for that future, and
20	position of children and adolescents and we'll	20	basic things like finance, they spent money that
21	come back to that with Professor Christie	21	had been given, that travelled the world, had
22	tomorrow, but can I ask you about what might be	22	made decisions based on the fact that their life
23	the particular psychological psychosocial	23	would not be long. I am very aware of the
24	challenges for those who have survived, who have	24	amount of conversations I've had with people who
25	aged with infection, who didn't expect to	25	would discuss that point and say, "I wasn't
	155		156
1	ready for that. It was really confusing, really	1	that in the broader setting of their lives, and
2	confronting to be able to have" or have to	2	the need for secrecy, the often distortions of
3	need to stand up and go, "Okay, you mean I'm now	3	communication that went on, somebody knew and
4	going to to live? Then I need to make some more	4	somebody else didn't know, and so on.
5	major adjustments to the life in front of me,"	5	So I think apart from, you know, the very
6	and had many wasted opportunities because they	6	serious impact of having a child with a life
7	did not believe they were going to live.	7	threatening condition, I think some of the sort
8	MS RICHARDS: Can I ask you about the psychological	8	of distortions in communication that almost
9	impact of everything you've described on family	9	I don't even want to call them games, but the
10	members, partners, parents, children, siblings	10	things people had to do just to keep their heads
11	and others.	11	above water, not telling people and a lot of
	Many of the those who were affected	12	avoidance went on, both by people who were
12	Many of the those who were affected through the infection of a loved one, who have	12 13	avoidance went on, both by people who were you know, avoidance in the sense of "Don't talk
12 13	•		
12 13 14	through the infection of a loved one, who have	13	you know, avoidance in the sense of "Don't talk
12 13 14 15	through the infection of a loved one, who have given evidence to the inquiry, have described	13 14	you know, avoidance in the sense of "Don't talk about it, don't deal with it". The avoidance
12 13 14 15	through the infection of a loved one, who have given evidence to the inquiry, have described psychological and emotional symptoms similar to	13 14 15	you know, avoidance in the sense of "Don't talk about it, don't deal with it". The avoidance seemed to be a very common response, and a very
12 13 14 15 16	through the infection of a loved one, who have given evidence to the inquiry, have described psychological and emotional symptoms similar to those described by the person who was infected.	13 14 15 16	you know, avoidance in the sense of "Don't talk about it, don't deal with it". The avoidance seemed to be a very common response, and a very understandable response, because, you know, why
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