
DOI: [10.1097/TXD.0000000000000581](http://dx.doi.org/10.1097/TXD.0000000000000581)

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Well, I Wouldn’t be Any Worse Off, Would I, Than I am Now? A Qualitative Study of Decision-Making, Hopes, and Realities of Adults With Type 1 Diabetes Undergoing Islet Cell Transplantation

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Background. For selected individuals with type 1 diabetes, pancreatic islet transplantation (IT) prevents recurrent severe hypoglycemia and optimizes glycemia, although ongoing systemic immunosuppression is needed. Our aim was to explore candidates and recipients’ expectations of transplantation, their experience of being on the waiting list, and (for recipients) the procedure and life posttransplant. Methods. Cross-sectional qualitative research design using semistructured interviews with 16 adults (8 pretransplant, 8 posttransplant; from 4 UK centers (n = 13) and 1 Canadian center (n = 3)). Interviews were audio-recorded, transcribed, and underwent inductive thematic analysis. Results. Interviewees were aged (mean ± SD) 52 ± 10 years (range, 30-64); duration of diabetes, 36 ± 9 years (range, 21-56); 12 (75%) were women. Narrative accounts centered on expectations, hopes, and realities; decision-making; waiting and uncertainty; the procedure, hospital stay, and follow-up. Expected benefits included fewer severe hypoglycemic episodes, reduced need for insulin, preventing onset/progression of complications and improved psychological well-being. These were realized for most, at least in the short term. Most interviewees described well-informed, shared decision-making with clinicians and family, and managing their expectations. Although life “on the list” could be stressful, and immunosuppressant side effects were severe, interviewees reported “no regrets.” Posttransplant, interviewees experienced increased confidence, through freedom from hypoglycemia and regained glycemic control, which tempered any disappointment about continued reliance on insulin. Most viewed their transplant as a success, though several reflected upon setbacks and hidden hopes for becoming “insulin-free.” Conclusions. Independently undertaken interviews demonstrated realistic and balanced expectations of IT and indicate how to optimize the process and support for future IT candidates.

(Transplantation Direct 2016;2: e72; doi: 10.1097/TXD.0000000000000581. Published online 21 April 2016.)
The second was also a mixed methods study, and demonstrated both the positive impact of IT on personal control over social life situation, and the recipients’ experience of IT as worthwhile. To our knowledge, no study to date has explored expectations and the process of undergoing IT from the individual’s perspective. Thus, our aim was to explore the expectations of people undergoing IT, how they weighed up advantages and disadvantages during the decision-making process, their experiences of being on the waiting list, the procedure itself and life posttransplant.

MATERIALS AND METHODS

Participants

All IT candidates and recipients at 4 UK transplant centers (King’s, Manchester, Newcastle, and Oxford) were invited to participate. Strict selection criteria for the UK IT program included: 2 episodes or more of severe hypoglycemia requiring third-party intervention over 2 years and impaired hypoglycemia awareness (Clarke questionnaire score ≥4), despite optimized conventional diabetes therapy. Risks and benefits of the procedure including alternative interventions were discussed with all potential transplant candidates, and all were assessed by a clinical psychologist before transplantation, with the main aim of identifying those who might require additional posttransplant support.

Due to the small numbers of UK transplants conducted at that time, and to gain an international perspective, a purposive sample of 6 participants was recruited from Edmonton, Canada: 3 pretransplant and 3 posttransplant, considered by their clinicians to have experiences that would enrich our study. Criteria for accessing the Canadian IT program were similar to the United Kingdom.

Posttransplant, immunosuppression therapy comprised tacrolimus/ sirolimus or tacrolimus/mycophenolate mofetil.

Interview Schedule

Using a semistructured interview schedule, we invited participants to explore their expectations and experiences in response to the following open questions: What did your doctor tell you to expect from your transplant? How much were you involved in deciding whether or not to have this sort of transplant? How do you feel about your transplant now? Posttransplant respondents were also asked: How satisfied are you with your transplant? Has the transplant met your expectations? If so, in what ways? Can you tell me about the immunosuppression drugs that you have to take?

Procedure

The UK Medical Research Ethics Committee gave ethical approval, with site-specific approvals at each UK center; the University of Alberta Health Research Ethics Board–approved interviews at Edmonton. Each participant provided informed written consent.

In the United Kingdom, interviews were conducted in the diabetes centers by 2 psychologists (rotating between J.S., A.W., and M.D.R.), all experienced in diabetes research and independent of the transplant teams. Two of the 13 participants were interviewed at home, accompanied by their husbands. For logistical reasons, interviews with participants based in Edmonton were conducted by telephone or “Skype.” Interviews typically lasted 90 minutes though
half of that time explored quality of life (not presented here). All interviews were digitally audio-recorded and transcribed, though 3 ‘Skype’ recordings were of poor quality, so those participants’ data were subsequently excluded.

**Analysis**

We used inductive thematic analysis and extracted quotes to illustrate themes. In contrast to deductive thematic analysis, where hypotheses are imposed upon the data, in inductive thematic analysis, the data are relied upon for generating the structure of the findings. This is the ideal approach when exploring new topics. Inductive thematic analysis involves several phases: familiarization with the data, generating initial codes, searching for themes among codes, defining and reviewing themes to produce a final structure. A “theme” constitutes a pattern of explanation given by more than 1 interviewee. Thematic analysis has been described as a tool that underpins several qualitative research methods, such as grounded theory and interpretative phenomenological analysis but is free from the theoretical constraints and assumptions imposed by those methods. More recently, thematic analysis has been considered as a valuable method in its own right.

In the first stage, between interviews, 1 psychologist (M.D. R.) read each transcript and proposed themes for each participant, making notes about codings. These were discussed by all 3 psychologists. In the second stage, 1 psychologist (A.W.) read repeatedly all transcripts and notes, together constituting the data corpus. At the end of thematic analysis, after repeated reading of the data corpus, no new themes were emerging. In the final stage, further checks (by J.S. and M.D.R.) confirmed no additional recurring themes.

An anonymized coding system—identity number (X), sex (M/F), transplant status (pre/post), country (UK/Canada)—is used to identify the source of each quote (in parentheses after each quote). Study center was excluded to preserve interviewee anonymity. Because most physicians were male and most transplant co-ordinators female, participants’ quotes were modified so that all physicians are referred to as males and all co-ordinators as females. Thus, professionals’ anonymity is preserved in the case of exceptions to this rule.

**RESULTS**

Table 1 summarizes the characteristics of the 16 participants (13 from the United Kingdom, 3 from Canada). Twelve were women, all were white, and 13 were married/cohabiting. The mean age of interviewees was 52 years, of whom most had received at least two separate infusions of hypoglycemia, which limited their quality of life; they referred to wanting “better control” or “greater stability” of blood glucose. Food and insulin impacted unpredictably on their blood glucose levels pre-IT. Post-IT, severe hypoglycemia was reduced or eliminated, at least short term, and awareness of hypoglycemia regained. However, the benefits did not endure for everyone. The earliest recipient interviewed had received 3 infusions but was experiencing frequent hypoglycemia again.

**Fewer “Hypos”, Better Awareness and “Better Control”**

Candidates were clear about why they wanted the transplant. Their reasons included recurrent unpredictable severe hypoglycemia (“hypos” or “lows”) and impaired awareness of hypoglycemia, which limited their quality of life; they referred to wanting “better control” or “greater stability” of blood glucose. Food and insulin impacted unpredictably on their blood glucose levels pre-IT. Post-IT, severe hypoglycemia was reduced or eliminated, at least short term, and awareness of hypoglycemia regained. However, the benefits did not endure for everyone. The earliest recipient interviewed had received 3 infusions but was experiencing frequent hypoglycemia again.

**Reduced Need for Injected Insulin: “No Guarantees”**

Interviewees were accustomed to injecting insulin, and most did not object to continuing to do so. Becoming insulin-independent was a pre-IT hope often tempered by realism and “no guarantee” clauses, which endured post-IT. Nonetheless, some retained hidden hopes to become insulin-independent and others had high expectations of “not having hypos, not being on insulin, being able to eat more or less what I wanted… everything about life would be better” (12FpostUK). The idea of the transplant “working” was tied up with hopes of insulin-independence. Even a small chance of an insulin-free period, no matter how short, went into the decision-making equation, even if it was then rejected as unrealistic.

**Post-IT, some had been temporarily insulin-free, although all were using insulin pumps or injections when interviewed. They could be “disappointed” or “upset,” particularly if denied another transplant, yet spoke of support from the**
transplant team and how they reframed their disappointment to focus on outcomes achieved, for example, fewer severe hypoglycaemic events, blood glucose stability.

“Not Being Diabetic”
Interviewees expressed the hope that having IT would mean they no longer had diabetes: “I’d be non-diabetic for a time... I would welcome the break... I’m very aware that at best I might get 5 years, and then ...be diabetic again, but...all the rest of it would be marvellous” (2FpreUK). Indeed, 1 posttransplant interviewee had declared to the UK Driving and Vehicle Licensing Authority that she no longer had diabetes rather than the diabetes controlling me” (9FpostUK)

Preventing or Halting Complications
Interviewees mentioned that pre-IT, they had hoped the transplant would have the benefit of preventing or halting

TABLE 2

<table>
<thead>
<tr>
<th>Sub-theme</th>
<th>Candidates’ (pretransplant) reports</th>
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<tbody>
<tr>
<td>Fewer ‘hypos’, better awareness and ‘better control’</td>
<td>“Not having the feeling of the hypo, the warning of the hypo, that is worrying... It’ll stop the hypos. I think... not having hypos is worth everything else, it really is” (3FpreUK)</td>
<td>“My life was made very unpleasant by hypos... I didn’t realise how appalling until it stopped. I couldn’t quite believe the difference... For the first time ever, I feel in control of the diabetes rather than the diabetes controlling me” (9FpostUK)</td>
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<td>“The hypos seem to be considerably less in people that have had them [transplants]. In which case, that would help me” (5FpreUK)</td>
<td>The last major hypo I had was the day before my first transplant” (13MpostUK)</td>
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<tr>
<td>Reduced need for injected insulin: “no guarantees”</td>
<td>“[The] transplant will not guarantee that I’ll be insulin-free and, you see, that doesn’t bother me” (3FpreUK)</td>
<td>“[I] was told that there was a possibility that I would come off injecting. I thought, ‘I’m not going down that line’. I just thought, ‘No, I think not somehow... and I think I went along that route because I just think it’s a more rational route” (9FpostUK)</td>
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<tr>
<td>Serious risks to immediate and long-term health</td>
<td>Swollen ankles, mouth ulcers, impaired liver function, impaired kidney function, headaches, tiredness... One wonders why I want to have it in the first place. But he is preparing me with statistics” (2FpreUK)</td>
<td>“Yeah, I was upset. But I had plenty of support [from named doctors]” (11FpostUK)</td>
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<td>“There’s 70% more chance of getting cancer, kidney problems, but they’ve not had that happen so far... and everyone’s got a chance of getting these things. I worried at first but not now... I figured other people had had it and they’re taking the drugs... I don’t dwell on it because if it’s going to happen, it’s going to happen” (8FpreUK)</td>
<td>I wasn’t even told that I would probably be able to come off insulin. I asked the question and they said, ‘It’s unlikely’. Now, they did it deliberately probably. They might have hoped that it could but they were certainly not intending that it shouldn’t happen.” (Posttransplant: “I was incredibly disappointed and then I thought, this is daft, why are you depressed?... What am I worried about? No major hypox, stabilisation of complications, they’re the most important things” (13MpostUK)</td>
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<td>“I must have some lucky guardian angel because, you know, I didn’t get anything, apart from a couple of colds, really it was luck I think, you know” (9FpostUK)</td>
<td>I think I was just disappointed. I was, you know, wanting the ability to stay off insulin but, again, if I still had the [blood glucose] control, taking needles wasn’t the problem” (16FpostCan)</td>
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<td>“I kept saying ‘I won’t get mouth ulcers. I won’t get anything’, and I didn’t” (12FpostUK)</td>
<td>“I kept saying ‘I won’t get mouth ulcers. I won’t get anything’, and I didn’t” (12FpostUK)</td>
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<td>“The only risk actually was from the drugs; the risk of the process itself is so minimal... and when I tried to get them to tell me, you know, out of 10 what are the risks... it was difficult for them, and I shouldn’t perhaps have tried to force it. But take for instance, the risk of developing skin cancer or remaining on anti-rejection drugs, um they can’t say that it’s you know 10% more than it was before or 20% more. I mean it’s ridiculous. All they can say is it’s an added risk, a real risk” (13MpostUK)</td>
<td>“The first lot of tablets, they were really making my mouth sore... very loose bowels, which is very, quite daunting, especially when you’re working” (14FpostUK)</td>
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<td>“They switched my immunosuppressants quite a few times... I had intense side effects” (16FpostUK)</td>
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progression of complications, particularly retinopathy. In the single case of islet after kidney, the interviewee indicated that the clinician was hoping “to protect my kidney for as long as possible”. Posttransplant, the only person who mentioned complications said that his had stabilized.

Benefits for Psychological Well-Being

The hopes of almost all IT recipients were realised, initially at least, with noticeably improved psychological well-being and energy: “I actually had energy… I didn’t feel so irritable” (9FpostUK). Another commented, “not embarrassing yourself at work or [with] friends, you know. That always bothered me… I don’t have any problems with low right now. So, yes, more confidence and less worries” (15MpostCan).

Serious Risks to Immediate and Long-Term Health

Overall, risks were less clearly recalled and described than benefits and were often considered unlikely. For people several years post-IT, the risks seemed to have passed. Interviewees were aware of, and some were fatalistic about, the possibility of immunosuppressant side effects. After a number of straightforward infusions, interviewees tended to place more emphasis on risk of side effects than on the procedure. Side effects ranged from severe, “I lost the end of my tongue actually from a terrible ulcer” (10FpostUK), to minor, “a couple of infections” or none, in which case, participants described themselves as “lucky” or having willpower.

| TABLE 3. |
| Decision-making pre-transplant and dealing with risks posttransplant |

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<thead>
<tr>
<th>Sub-theme</th>
<th>Candidates’ (pretransplant) reports</th>
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<tbody>
<tr>
<td>Informed decision-making</td>
<td>“Yes, I realise that the anti-reduction, rejection drugs can be nasty, um, and I think they stressed that, possibly over stressed it, but I can see why they’ve done it because you don’t want somebody afterwards saying, ‘Well, you didn’t tell me about this and you didn’t tell me’… they’ve been very, very open about the things that could happen” (3FpreUK)</td>
<td>“We made a decision as a family… to say yes this should improve things for all of us” (10FpostUK)</td>
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<td>“Realistically, at the end of the day, it’s always going to be my choice, isn’t it?” (4MpostUK)</td>
<td>“It was very much a joint decision to my wife and me” (13MpostUK)</td>
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<td>“It sounds exciting - anything that helps hopefully is going to be good, but you need to be more informed about everything don’t you? But I’m willing to learn everything I can” (5FpostUK)</td>
<td>“Together (my endocrinologist and myself) we made the decision. Then... he gave me what I needed to know and for me to go home to my family” (16FpostCan)</td>
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<tr>
<td>Involving others in the decision</td>
<td>“I speak to my husband, and... I don’t know if I would involve children. I don’t think it’s probably fair... they’re concerned, but they also want the best for me” (2FpostUK)</td>
<td>“We were very much a joint decision to my wife and me” (3FpostCan)</td>
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<td>“Well I spoke to my son and he said, ‘There’s risks with everything’... He’s saying, ‘Go for it, just go for it!’ and I said, ‘Well, yes, I do want to, but I’m only trying to think of what the minus is on it.’ You’d be a fool not to, wouldn’t you really? You have to think of that but on the whole... it sounds good” (5FpostUK)</td>
<td>“When I tried to get them to tell me, you know, ‘Out of 10, what are the risks, developing this or that?’, it was difficult for them” (13MpostUK)</td>
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<tr>
<td>Balancing risks against benefits</td>
<td>“If you thought that would happen, you wouldn’t have it in the first place. You hope it won’t happen to you” (2FpostUK)</td>
<td>“There wasn’t really a lot of information, um, on the procedures... it’s been done, you know, four times before” (14FpostUK)</td>
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<td>“It’s like everything, what affects one won’t affect another. Well, anything can go wrong, but... I don’t think anything is without its risks, um, and I do believe if it’s going to happen, it’s going to happen, whatever it is” (3FpostUK)</td>
<td>“Yes, there are obviously risks. There are risks to everything. You walk across the road it’s a risk, isn’t it?” (8FpostUK)</td>
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<td>“I don’t want to get cancer but... I can’t let it stop me” (8FpostUK)</td>
<td>“When I tried to get them to tell me, you know, ‘Out of 10, what are the risks, developing this or that?’, it was difficult for them” (13MpostUK)</td>
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happen was that the procedure did not “work,” and concluding “Well, I wouldn't be any worse off, would I, than I am now?” (5FpreUK). Post-IT, after a number of straightforward infusions, recipients emphasised the risk of the immunosuppression side-effects rather than the procedure.

With regard to ongoing immunosuppression required post-IT, the reality was that some experienced health problems. However, none of the candidates or recipients spoke of it in terms of swapping 1 treatment (insulin) for another (immunosuppression). Even those who had experienced several setbacks reflected that there was no permanent damage: “I tried something and it didn’t work and that’s pretty much it” (16FpostCan).

**Waiting and Uncertainty**

**Pre-transplant Investigations: Not Building up Your Hopes**

During pre-IT investigations, participants did not want to build up their hopes. The time commitment for pre-IT investigations was considerable and this extended period of uncertainty could be stressful (Table 4).

**On the Waiting List: “Is the Phone Alright?”**

The wait for a suitable donor was often lengthy, stressful and limited interviewees’ ability or willingness to travel. Post-IT interviewees described this as the most difficult part, becoming increasingly anxious when there was no call from the centre.

**False Alarms**

When the call came, the islet preparation could prove unsuitable. Participants experienced at least one false alarm, which could be “upsetting” or depressing.

**After Transplant: “It’s in the Lap of the Gods”**

Even when the transplanted islets produced insulin, there remained uncertainty about the future, whether they would remain insulin-independent and whether or not subsequent transplants would be needed.

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**TABLE 4.**

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<tr>
<td>Pretransplant investigations: not building up your hopes</td>
<td>“… if I am the recipient, um, because I don’t think it does to build your hopes up too much. I shall be very, very disappointed if I’m not. But if I’m not, ‘so be it, you know” (3FpreUK)</td>
<td>“I think it would be nice to sort of hurry up the investigations and stuff, because it takes like 3 months, and you’re wondering… if you’re going to be listed and stuff… I didn’t realize it would take so long [Then, waiting for 2nd infusion] They seem to be more on top of it now and said, you know, ‘We’ll get you in for your bloods Monday because we don’t want to keep you waiting’” (11FpostUK)</td>
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<td>“The decision isn’t really made insofar as, there’s still work that’s got to be done, um, different tests to do. I could fail at any one of those hurdles” (6FpreUK)</td>
<td>“They tried to make it as simple as possible but there are probably 10 different times you had to come in for a certain test or something like that. So, I would estimate about 10 times and then the wait for the transplant” (16FpostCan)</td>
<td></td>
</tr>
<tr>
<td>On the waiting list: ‘Is the phone alright?’</td>
<td>“[Waiting …] was almost the worst bit because, you’re completely on edge the whole time and every phone call at a weird time you think, ‘Oh God, that could be it.’” (10FpostUK)</td>
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<td>“Every time the phone went I expected it to be the hospital… I missed my son’s graduation… I had to cancel holidays” (13MpostUK)</td>
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<td>“I waited about…] 6 to 8 weeks I think. It seemed like quite a long time at the time. I was like, ‘No-one’s ringing, no-one’s ringing! Is the phone alright?’” (14FpostUK)</td>
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<td>False alarms</td>
<td>“I got one… a call, just before Christmas, to go up. I was up there the whole day and it wasn’t suitable for me at the end of the day, so that was quite an upsetting sort of issue!” (11FpostUK)</td>
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<td>“You get all hyped up, you’re there and ready to go and the next thing, you’re all fixed up with your IVs and they say, sorry, you’re not getting it this time, the cells aren’t good enough, there weren’t enough cells and it was kind of a low period for a while there” (15MpostCan)</td>
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<td>“They may want to do a third transplant, they might not, I don’t know, because I don’t know what’s going to happen to these hormones… I mean, I don’t know… it’s just, it’s in the lap of the gods” (9FpostUK)</td>
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**The Procedure, Hospital Stay and Follow-Up: “No Regrets”**

Pretransplant interviewees thought they had a clear understanding of the procedure and described it in their own words (Table 5).

When the procedure went well, it was mostly described as quick, pain-free and unproblematic: “it took half an hour, and that was extraordinary” (9FpostUK). However, some experienced “painful” complications.

Recovery was usually swift but when the procedure was not straightforward, hospital stays were longer than expected. In the United Kingdom, because it was a novel
procedure, the hospital stay was routinely longer than that in Edmonton. Posttransplant check-ups were a considerable commitment, and particularly burdensome if living far away from the centre. Some said they had not realized “how extensive it would be” (16FpostCan).

Balancing the outcomes against immunosuppressant side effects, the procedure and its eventual outcome, as well as the time commitment, most reported “no regrets” about “going for it”.

**DISCUSSION**

This is the first in-depth qualitative study investigating the expectations and experiences of adults with T1DM undergoing IT in the United Kingdom, without which meaningful quantitative evaluation of IT is not possible. Although several quantitative investigations have evaluated impact on health and well-being,

our study represents the first to explore expectations, hopes, decision-making, and outcomes in the recipients’ own words.

Although none of the post-IT participants were insulin-free at the time of interview, insulin independence has never been promoted actively as a goal of the UK islet transplant program. All interviewees reported severe recurrent and unpredictable hypoglycemia pretransplant and most expected or hoped for modest improvements afterwards. Consistent with published quantitative findings,

this study confirmed that, posttransplant, interviewees reported being pleased with various combinations of improved glycemia, regained awareness of hypoglycemia, and, most commonly, reduced frequency or severity of hypoglycemia. This had “life-changing” psychological benefits, improving well-being, confidence, and quality of life.

Overall, in contrast to the post-IT participants (early recipients), transplant candidates reported realistic expectations, especially concerning “coming off insulin.” Physicians counsel about this possibility pretransplant. Although transplant teams largely managed expectations appropriately, many interviewees retained hidden hopes (reportedly shared by some physicians) of being among the minority to remain insulin-free at 5 years. This natural tendency toward optimistic bias needs to be recognized and managed by transplant teams. Furthermore, recipients need support to develop strategies to cope with disappointment at any stage, including not being “suitable,” lengthy wait for donor organ, “false alarms,” graft rejection, or function loss. Even though transplant teams are now careful not to frame IT as a potential cure, the lay belief endures, meaning that extra care is needed to ensure that all professionals give the same cautious message. It has been acknowledged elsewhere that individuals’ goals and expectations impact on their perceptions of IT success.10

Whether or not a period of insulin independence had been experienced, coping with any disappointment when insulin became necessary was addressed by reframing the benefits to focus on blood glucose stability. A determination to remain positive, combined with the negative consequences of transplantation, has been noted elsewhere.21,22 Whereas kidney transplantation is life-saving and success is dichotomous (ie, independence from dialysis or not), IT is life-changing and offers several potential benefits. Thus, whereas coping with disappointment and avoiding depression would be difficult for kidney transplant recipients,21 it may be easier for IT recipients who can focus on other benefits.

Coping strategies were also apparent in participants' descriptions of how they dealt with the uncertainty of life “on the list,” with emphasis placed on luck, fate, chance, and willpower. Such observations have been made, where transplant recipients used external loci of control to manage uncertainties and organ rejection.23 Evidence from kidney transplant research suggests that belief in chance or fate may be a reasonable coping mechanism, because those with greater perceptions of internal or personal control were most depressed when the transplant failed.24,25

Given the realities of the transplant process, uncertainty for those on the waiting list is unavoidable. Candidates need to be prepared for the significant time commitment involved in medical investigations. Pre-IT and post-IT investigations need to be conducted efficiently, with appreciation that each

**TABLE 5.**

The procedure, hospital stay, and follow-up

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<tr>
<td>‘No regrets’</td>
<td>“The islets will be injected into my liver… I think, is it the portal bit that goes into the liver? Anyway… I know it’s going to be done under local [anaesthetic]… I don’t know how long it’s going to take, um, I’m presuming it’s going to be drip fed in at a controlled rate… and I know it’s going to be done in the X-ray department so that they can see… I think [the cells] sit in the liver and they work in the liver” (6FpreUK)</td>
<td>“The second procedure took an hour and a half, um, because it was more difficult to get the actual tube into the portal vein… and I thought ‘this could be a new form of torture’ … I felt pretty fragile for a day and then I began to perk up no end” (10FpostUK)</td>
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<td>“It was really painful to breathe, because something had happened with my lungs, um, so um, I was on morphine and things that week, but the ones after that were absolutely fine” (16FpostUK)</td>
<td>“It didn’t hurt me; it wasn’t painful; there was nothing unpleasant about any of it” (12FpostUK)</td>
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<td>“I had some problems right after the transplant. I was in hospital for a week. It wasn’t a happy time for me… I thought I’d just have the transplant and be home in a day or so” [Post-IT check-ups] “I used to go three times a week two weeks a day… It was very restrictive. First thing in the morning at 7 o’clock… then come back two hours later, and they did that for three months” (15MpostCan)</td>
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individual balances this against personal commitments, including family and employment. A successful transplant program needs to prepare participants fully for the risks involved—with sensitivity to how much information potential candidates may or may not want before their “suitability” as a candidate is confirmed—both to physical health and psychological well-being, and to detect problems and provide support at any stage. Timely and efficient investigations, psychological assessments and a responsive clinical team, who provide consistent messages, are key ingredients for ensuring best possible care for IT recipients and their families during this time of high hopes and unavoidable uncertainties. Our findings have informed the published Diabetes UK patient guide, particularly ensuring that information about risks of comorbidities and anticipated frequency of hospital attendances are explicit.

Unsurprisingly, the decision to join the transplant waiting list was made by the individual with support from, and information sharing with, their clinical team. Family involvement was actively encouraged but the individual decided when to involve them and often needed to counter the families’ enthusiasm by advocating information-seeking about the risks and realities to reach a rational decision.

Our study has several limitations, particularly relating to the sample. The UK sample size was small but included all IT recipients and candidates (those on the waiting list) at the time; the Canadian purposive sample was added to maximize confidence that UK experiences were similar to those in a more established program. The selective and subjective sampling of Canadian participants and the fact that several recordings were of poor quality is a limitation. However, we noted considerable heterogeneity of experience among the Canadian participants, and their inclusion was only ever intended to augment the sample size and validate the UK experience, this being the main focus of the study. This remains the largest qualitative study of IT conducted to date. Qualitative studies cannot claim to be representative but after repeated reading of the data corpus, no new themes emerged, and we were satisfied that data saturation had been achieved. We had a unique opportunity to understand the experiences of IT recipients and candidates at an early stage of the UK program. There was remarkable commonality between the accounts of United Kingdom and Canadian participants. This may be explained by the fact that there is still a small number of IT centers, with close collaboration and intercenter support internationally. Although only half the participants were posttransplant, which limits the findings about posttransplant status, there was strength in this approach because the experience of being on the waiting list was current for half the participants, not relying on retrospection or subject to potential reframing. The diversity of centers (4 United Kingdom and 1 Canada) can be considered a strength, whereas the unintentional preponderance of whites and women raises some concerns for generalizability. The IT program itself has a clear bias toward female recipients, given the need for relatively low body weight and modest insulin requirements. It is noteworthy that the only other qualitative study also includes more women than men. It is possible that IT recipients from other ethnic groups may be more trusting and less willing to ask questions, particularly if that might be viewed as challenging authority.

Another limitation is that many early recipients had achieved “celebrity status” in the diabetes world and, clearly, all were grateful to their clinicians for the opportunity to be among the first to receive an IT. Their many press interviews may have led to some rehearsed responses and a certain reticence to highlight any negative aspects of IT. To maximize the likelihood of unfiltered and balanced feedback, we used 3 strategies: (1) the psychologists who conducted the interviews were independent of the clinical centers and direct care of the transplant recipients; (2) participants were assured of anonymity, with any identifying words removed from the quotes used; (3) we tapped into the participants’ altruistic tendencies and support for research by assuring them that we were interested in any experiences/comments that would help the centers to improve the islet transplant experience for future recipients. Before joining the IT waiting list, prospective candidates were screened for psychological stability, meaning that extreme emotional reactions or unrealistic expectations were unlikely. Further research with a larger sample from the more established UK program, and considering the longer-term outcomes, is now warranted to corroborate or update these findings, especially since the introduction of the new Diabetes UK patient guide to IT.

This independently undertaken qualitative study sought to capture the breadth and depth of experience of those undergoing IT. It provides insights into both common and individualistic expectations and experiences, offering clinicians the opportunity to learn how to optimize the process, assessment, and support for future IT candidates. The study also provides a person-centered evidence base for developing and selecting robust questionnaires for quantitative assessment of patients’ expectations, experiences and satisfaction with IT, alongside biomedical outcomes.

ACKNOWLEDGMENTS

The authors thank the people awaiting IT or having received IT who participated in this study, for generously sharing their time and insights.

The authors also thank the transplant co-ordinators and other center staff who helped with participant recruitment: Edmonton: Parastoo Dinyari, Sharleen Imes, Lana Toth, Andrew Malcolm, Angela Koh; King’s: Andrew Pernet; Manchester: Sarah Pendlebury with infrastructural support from the Manchester NIHR Biomedical Research Centre; Newcastle: Julie Wardle, Charlotte Gordon; Oxford: Anne Brownson, Neil Walker.

The authors thank representatives from all clinical centers within the UK Islet Transplant Consortium who supported, reviewed, and approved the work reported: Richard Smith (Bristol); Shareen Forbes (Edinburgh); Pratik Choudhary (King’s College London); Gareth Jones and Miranda Rosenthal (Royal Free London).

REFERENCES