Specimen Collection for Induced Pluripotent Stem Cell Research: Harmonizing the Approach to Informed Consent

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ABSTRACT

Induced pluripotent stem cells (iPSCs) have elicited excitement in both the scientific and ethics communities for their potential to advance basic and translational research. They have been hailed as an alternative to derivation from embryos that provides a virtually unlimited source of pluripotent stem cells for research and therapeutic applications. However, research with iPSCs is ethically complex, uniquely encompassing the concerns associated with genomics, immortalized cell lines, transplantation, human reproduction, and biobanking. Prospective donation of tissue specimens for iPSC research thus requires an approach to informed consent that is constructed for this context. Even in the nascent stages of this field, approaches to informed consent have been variable in ways that threaten the simultaneous goals of protecting donors and safeguarding future research and translation, and investigators are seeking guidance. We address this need by providing concrete recommendations for informed consent that balance the perspectives of a variety of stakeholders. Our work combines analysis of consent form language collected from investigators worldwide with a conceptual balancing of normative ethical concerns, policy precedents, and scientific realities. Our framework asks people to consent prospectively to a broad umbrella of foreseeable research, including future therapeutic applications, with recontact possible in limited circumstances. We argue that the long-term goals of regenerative medicine, interest in sharing iPSC lines, and uncertain landscape of future research all would be served by a framework of ongoing communication with donors. Our approach balances the goals of iPSC and regenerative medicine researchers with the interests of individual research participants.

INTRODUCTION

Human induced pluripotent stem cell (iPSC) research activity has seen a monumental expansion since the discovery of somatic cell reprogramming in 2006 [1, 2]. These cells have been hailed as a panacea for basic and translational medical research; the potential of this technology ranges from disease modeling and genetic analysis of otherwise inaccessible tissues to candidate drug screening to immunologically matched cell transplant therapies and novel infertility treatments (Table 1) [3–51]. Although recent evidence has tempered the hope that translating these technologies toward new therapies will be easy [52–64], there is great interest in using iPSC lines to advance translational goals [65]. A broad range of human tissue types are currently being procured to facilitate the generation of iPSC lines [13, 23–26]; ensuring that the prospective collection of specimens for this research proceeds with appropriate informed consent is thus a central objective [66, 67].

Prospective donation of tissue specimens for iPSC research requires a nuanced approach to informed consent. Many of the salient features of iPSC research are no different from research involving other pluripotent stem cells, immortalized cell lines, human biospecimens, genetic and genomic analyses, reproductive research, and transplantation [67]. However, although the individual issues associated with these cells are not new, iPSC research uniquely encompasses them all. iPSCs are immortal, creating longitudinal challenges related to withdrawal, confidentiality, and sharing. They are pluripotent, and the range of possible uses of these cells is ever-growing. They are likely to undergo genomic analyses for intensive characterization and disease screening, marrying these cells to the ethical concerns of genomics. Furthermore, because these cells can be sourced from almost any kind of specimen and can be used for potentially limitless rounds of derivation and paths of differentiation, there is the potential for unprecedented flexibility in derivation, sharing,
Table 1. iPSCs: the state of the science

| Fundamentals | Current research | Potential | Challenges |
|--------------|-----------------|-----------|------------|
| **Seminal studies** | Takahashi et al. [3] | Basic | Basic | Basic |
| | Yu et al. [4] | Stem cell biology [9] | Biological models and mechanisms of disease [37–42] | Genomic instability [52–55] |
| | | Reprogramming methods—retrovirus, lentivirus, adenovirus, Sendai virus, episomal plasmids, mRNA, proteins, and small molecules— the last 5 are nonintegrating and minimize risk of mutagenesis and other complications [10–18] | Elucidation of stem cell development and pluripotency | Epigenetic instability and memory (of original lineage) [56–58] |
| | Unique qualities | Optimizing reprogramming, genetic correction and modification, culture maintenance, and differentiation [19–22] | Genetic research | Reprogramming method (integration free approaches are ideal) |
| | | Use of different cell types as sources (cord blood, peripheral blood, marrow, fibroblasts, B cells, progenitors, etc.) to improve efficiency of reprogramming and subsequent differentiation [23–26] | Chimera research | Efficiency of reprogramming |
| | | Biomaterials approaches to improving culture/transplantation [27, 28] | Ability to access tissues otherwise difficult to obtain | Ease and reproducibility of culture |
| | | Transdifferentiation [29–31] | | Optimizing cell culture/niche conditions |
| | Translational | iPSC lines derived from patients with specific genetic variants (both Mendelian and complex) to be used to screen for small molecules to cure or reduce clinical effects | Translational | Engraftability |
| | | Development of cell transplants for conditions ranging from premature hair graying to Parkinson’s disease [32–36] | Cell and whole tissue assays to screen new pharmaceuticals for efficacy, safety, and toxicity [43] | Clinical grade production [63] |
| | | | Cell transplantation and regenerative medicine: patient-matched cell therapies to treat psychiatric and degenerative conditions, cancer, traumatic injury, and genetic disease (often in combination with other drugs or tissue-engineered scaffolds) [44–49] | Comparative |
| | | | Gametogenesis, assisted reproduction, infertility, and developmental research [50, 51] | Scientific and therapeutic benefit compared with embryonic stem cells (gold standard) [64] |
| | | | | Requires parallel hESC research to ensure studies are valid (thus, iPSCs do not completely eliminate the ethical and political hurdles); public misconception that iPSCs are a wholesale replacement for hESCs |
| | | | | Comparison with adult stem cells |

Abbreviations: hESC, human embryonic stem cell; iPSC, induced pluripotent stem cell.
and banking. The scientific potential of iPSCs and the future therapies they make possible are extraordinary. In addition, research with iPSCs is unique because of the drive to transition rapidly toward therapy, making translational goals fundamental to this research from the outset.

Although a clear set of research and translational goals for iPSC research has been articulated [65], the precise nature of future research is largely unknown at this time, and providing comprehensive information to prospective donors of specimens for iPSC research via the consent process is especially challenging. This is important, given data about public attitudes that suggests there may be hesitation to participate in the earliest stages of research and significant concerns about germ line cell derivation and reproductive applications, although there is general support for iPSC research with appropriate ethical and regulatory oversight [68]. The process of contributing specimens to iPSC research can open doors to significant applications and associated concerns that are not yet foreseeable by scientists, the magnitude of which are incongruous with the relatively facile donation process and are difficult to convey in a consent form.

We argue here that the long-term goals of regenerative medicine, broad interest in sharing iPSC lines among scientists, the largely unknown landscape of future research, and sensitivities associated with some potential uses all point to a consent process that comprehensively addresses these goals and concerns. We describe an approach that asks people to consent prospectively to a broad umbrella of foreseeable research, anticipating downstream issues and goals to the extent possible by including future therapeutic applications of iPSCs. We believe this is the best approach among multiple ethically acceptable avenues. It balances the goals of iPSC and regenerative medicine researchers with the interests of individual research participants. By granting that participants have longitudinal interests in the use of their specimens, and by providing narrowly defined mechanisms by which to exercise those interests, our approach encourages a framework of ongoing communication between researchers and participants. This approach will allow scientific progress to continue forward while ensuring that donors of specimens are respected when providing consent to such a broad scope of future research.

**BACKGROUND**

There is value in developing a consistent approach to informed consent across various research institutions in these still-early stages of regenerative medicine. Inadequate consent processes can both undermine the public’s trust in research with cell lines, as the HeLa cell case teaches us [69], and hinder future research with cell lines. Problematic variability has been documented in consent forms for research involving the collection, storage, and future use of biological specimens [70, 71]. Such variation has complicated the ability of researchers to use embryonic stem cell lines [72–74]. Anticipating future applications and associated regulatory requirements now, to the extent possible, will help balance the goals of protecting participants with maximizing the utility of iPSC lines in research. Although it is not possible to predict which iPSC lines will be the “next HeLa” and lead to important therapeutic discoveries, following careful consent procedures for all generated iPSC lines will both ensure ethical provenance of all cell lines as they traverse many potential research and distribution pathways (Fig. 1) and protect the ability to utilize rare and valuable discoveries associated with any of the cell lines down the road.

A variety of approaches to informed consent for research on specimens can be ethically justified [75, 76]. Some argue that research on specimens is well-suited to broad and open-ended consent approaches that involve a single interaction between the researcher and participant [75, 77, 78]. Policies that facilitate the systematic use of one-time consent for specimen research have been incorporated into proposed reforms to the U.S. human subjects regulations [79]. Such models assume that ongoing interaction with participants is neither necessary for the research nor desirable from the participant’s perspective, and studies suggest that many participants are indeed comfortable with a one-time approach to consent for research with their specimens [77, 80].

However, there remain concerns about broad, one-time consent: specifically, questions about whether and how prospective participants can give adequate informed consent if they cannot be informed at the time of donation about some of the possible ways that their specimens will be used in the future. Allowing a participant to give carte blanche permission to unbounded future research may be inadequate in the case of iPSCs, as it is reasonable to assume that most participants are not truly aware of what such a broad permission could entail [81]. Broad, open-ended approaches may deter a small number of participants from enrolling in research. Although this may be a reasonable trade-off for some categories of research, it would be problematic if those with acute or rare diseases, who have the most at stake and whose specimens are also likely valuable, are disproportionately deterred from participation by being required to sign on so broadly.

At the other end of the spectrum are proposals to construct consent forms narrowly, limiting them to study specifics that are known at the present time [75, 76, 82]. Such approaches would likely assume that it is premature to prognosticate about future therapeutic applications of iPSCs, at least in most cases. Although broad up-front consent might be allowable in a limited number of cases—for example, if scientists had the ability to determine that a specific cell line would be particularly useful—researchers would generally be required to recontact participants to get consent for each new project and application that uses their iPSCs. However, there are several problems with this approach. First, a requirement for repeated reconsent in most cases is inefficient and would hinder the ability to share and use samples and cell lines broadly. In addition, it is highly unlikely that researchers can predict at this time which cell lines will be clinically useful in the future; this is not currently a useful metric for choosing between broad versus narrow consent. The current state of scientific knowledge requires us to assume that all iPSC lines have potential future clinical utility, even if only a few lines will actually go on to be used in therapeutic applications. Clinical trials with human embryonic stem cells (hESCs) show that it is possible to make clinically compliant samples from research grade materials [63], and samples obtained through blood and cord blood banks and bone marrow registries are already collected in a clinical-grade manner [23]. Finally, we are skeptical that narrowly tailored consent approaches that require frequent recontact of participants substantively enhance protections or
respect for participants in iPSC research more so than approaches that include broader descriptions of iPSCs and their future applications [72].

On balance, these concerns about overly broad and overly narrow approaches to consent point to a process that provides accurate information about the broad aims of iPSC research and downstream goals, draws boundaries around the scope of the consent, and establishes an ongoing dialogue with participants that allows for reconsent in some cases. We believe such a middle ground exists.

**REGULATORY AND ETHICAL FRAMEWORKS**

Before delving into the content of consent forms, it is helpful to define the purpose of informed consent for iPSC research. The overarching goal is to provide sufficient baseline information that enables potential participants to decide whether to give permission for iPSC research to proceed with their specimens [83]. The consent process respects the autonomy of participants by giving them reasonable control over the use of their specimens. Informed consent also has the potential to establish a broader relationship between investigators and participants aimed at mutual benefit, trust, and education [84]. The degree of control that is granted to participants, the frequency of interactions between researchers and participants, and the level of detail about possible research applications are all dimensions of the consent process that need to be decided.

Tracing the development of the discourse surrounding iPSCs also helps frame the requirements for informed consent. Because iPSCs obviate the need for source blastocysts, they were initially hailed as ethically superior to hESCs [85–88]. Yet iPSCs...
are not wholly free from the encumbrances of prior ethical debates regarding hESCs [89–92]. Global objections to iPSC research related to its complicity with embryonic stem cell research (arising from the necessary symbiosis between the two research programs), its ability to alter our conceptions of human life, and its implications for human-animal chimeras have been raised [66, 93–102]. Although these questions continue to be debated, both publicly and privately funded research with iPSCs is proceeding with broad scientific, political, and public support and without the heavy regulation that has characterized research with hESCs [103, 104].

Only recently have iPSCs been recognized to have more complex ethical dimensions [67], presenting concrete ethical and logistical issues analogous to those with which biobanks, stem cell research oversight (SCRO) committees, institutional review boards (IRBs), and other bodies have previously struggled including terms of use, confidentiality, tracking, governance, and withdrawal [105–109]. There is also uncertainty about the role that the U.S. Food and Drug Administration (FDA) will play in regulating this research and translation [110–115]. Of particular concern for researchers involved in regenerative medicine is anticipating how FDA oversight and regulation will affect the technology transfer and translation of iPSC-related materials and therapies, what up-front and ongoing requirements will need to be satisfied [infectious disease testing, at a minimum], and what role, if any, informed consent will play. The informed consent process for research with iPSCs needs to take this complex ethical backdrop into account [75, 116, 117]. An emerging literature highlights the value of “tiered” approaches that allow participants to make choices [67], and describes traceability and withdrawal concerns as they apply to consent for stem cell banking [84]. Guidelines at all levels of oversight—international bodies and working groups [118–121], federal agencies [122, 123], state boards and foundations [124, 125], even institution-level IRBs and SCROs—are beginning to speak to the complexities of regulation and informed consent in these contexts by recommending new approaches, including specific provisions for human transplantation, recontact, stem cell banking, opt-out for return of results, tiered consent, and partial withdrawal. Our analysis builds upon this foundation.

**METHODS**

We developed a model consent template for the prospective creation of iPSC lines for research. This template was informed by (a) a conceptual analysis and normative balancing of issues associated with the various informed consent content domains, and (b) a content analysis of 25 iPSC-specific consent forms and previously approved language that were shared with us by investigators and administrators from a variety of U.S. and international institutions (Table 2). Our analysis incorporates a review of relevant literature and careful tailoring of various consent domains to the iPSC context. The example consent forms were reviewed for clarity as well as content and compared with each other to uncover qualitative patterns. In addition, our proposed template was informed by conversations with a variety of stakeholders, including investigators, bioethicists, IRB chairs, lawyers, research administrators, and those involved in federal and state-level stem cell research policy and regulation.

This project was a joint effort between the newly formed National Institutes of Health (NIH) Center for Regenerative Medicine and the NIH Clinical Center Department of Bioethics, with the goal of producing a consent template for prospective collection of fresh specimens to create iPSCs for research that would be useful both for our intramural investigators and the broader research community. The guidance provided here is meant to take advantage of a window of opportunity to harmonize the guidance provided to investigators and institutions, but of course is not a substitute for IRB approval of individual studies.

A summary of the provisions that were incorporated into our model consent template (supplemental online data) is provided in Tables 3 and 4 and discussed below.

**ANALYSIS OF CONSENT DOMAINS**

We evaluated the content domains of informed consent that are most relevant to iPSC research and concluded that the most salient ethical concerns arise from the scope of possible future research applications, both foreseeable and unknown, of iPSCs. Many potential uses—reproductive research and gamete generation, mixing of human and animal biological materials, pharmaceutical screening (separation between “research purposes” and commercial development for profit), transplantation into humans for regenerative medicine, and genetic sequencing and manipulation—are essential as both methods and paths for future exploration yet are potentially sensitive in nature and may reasonably elicit objection from prospective participants. These uses sort into three general categories: core foundational methodologies, regenerative medicine, and reproductive research. Furthermore, it can be assumed that there are potential uses of these cells that cannot yet be predicted, some of which may be sensitive. The consent process should provide sufficient information to allow participants to be aware of all of these possibilities.

We acknowledge that participants have an inherent stake in how their donated specimens are used, and thus have longitudinal interests in the research performed on the generated iPSC lines generated. This gives rise to several additional concerns that are appropriate to address in the informed consent process. These concerns can be sorted by the general principles and categories of informed consent. Although these categories are not necessarily unique to iPSC research, the discussion proceeds with an eye toward how they should be applied to that context.

**Confidentiality, Traceability, and Sustained Interaction**

A description of risks to confidentiality and measures that will be taken to minimize these risks are required elements of informed consent (as per 45 CFR 46.116) [126]. This information is important in part because it allows participants to make informed decisions about the acceptable amount of privacy risk.

Traceability (i.e., maintaining accessible coded identifiers rather than de-linking samples and associated data) may be beneficial in the case of pluripotent stem cell banking. A system of traceable specimens can facilitate ongoing communication, participation, and information exchange. A “sustained interaction,” so defined, would benefit the research enterprise by fostering public trust and streamlining the translation of research [84]. Traceability can also facilitate longitudinal data collection, create...
an avenue for return of research results, and enable participants to be informed of research outcomes [84].

Traceability does entail greater risks to confidentiality, requiring explicit language in the consent form about the protections being afforded. Although de-linking or “anonymization” significantly reduces risks to confidentiality, these measures may limit the scientific utility of the lines, as well as any associated benefits to participants. Furthermore, these approaches do not address the ethical problems associated with the inherent stake that participants have in the ongoing and future uses of their specimens [72], and limited empirical evidence shows that participants do not necessarily favor this approach [127]. A balance must be struck that takes into account scientific considerations, privacy concerns, obligations to maintain ongoing contact with participants, and practical administrative realities.

Return of Benefit and Information
Closely related is the idea of return of benefit through the provision of research results. Prospective participants are unlikely to benefit from iPSC research directly — either through a direct medical intervention or through future access to “personal” stem cells. However, there remain open questions about obligations to return either aggregate or individual research results, validity of individual results when obtained downstream from mutated or genetically altered cell lines, and methods to capture participants’ preferences regarding receipt of such results. Any plans for recontact and return of results, whether robust or limited, should be described up front in the consent process.

Commercial Product Development
Because iPSCs will potentially be used as drug screening models or direct transplant interventions, commercial profits may be possible. The potential for profits connected to iPSC research should be clearly expressed to participants, as this may affect a participant’s conception of his or her “donation.” Such disclosures are prompted largely by legal precedents [128] but are motivated by ethical considerations as well. Some argue that sponsorship, commercialization, and financial interests are relevant to a participant’s assessments of risks and benefits, and call for exhaustive disclosure of commercial aspects prior to enrollment. However, this must be weighed against the need for concise consent forms to better promote understanding, which is already quite challenging [129, 130]. The potential for commercialization may be relevant to participants’ decisions to allow their cells to be used therapeutically.

Withdrawal
A right to withdraw from participation in biomedical research is widely acknowledged in the human subjects research ethics literature [131]. How this right extends to donated specimens and pluripotent cell lines is unclear, however, particularly for those samples that will be stored and distributed widely. The extent of a right to withdraw is further complicated by questions about whether cell line derivatives still constitute human subjects research. There are many possible avenues for withdrawal that vary greatly in both practicality and adequacy: ceasing recontact, anonymization (destruction of identifiers), destruction of original specimens, ceasing distribution of any materials (original or derivative) and information, complete destruction of iPSC lines, and efforts to ask partner institutions to anonymize or cease use of the shared iPSC lines. The choice of an appropriate approach should take into account religious beliefs and cultural traditions that require specimens to be returned to participants [132].

Storage, Banking, and Exchange
Provisions for sharing and storage of iPSC lines may be of particular interest to participants, including whether these cells will stay with the primary investigator, will be shared nationally or internationally, and/or will be banked in a widely available cell repository. Institutions may vary in their governance of the use of these cells and in confidentiality protections, and restrictions promised by the initial investigator may not transfer when shared. Participants should be informed about oversight assurances and provisions for future uses, regardless of goal or location of distribution.

Recontact and Reconsent
Recontact permits evolving issues related to the research to be addressed with participants on an ongoing basis. Some have recommended recontact of participants in pursuit of reconsent if iPSCs derived from their specimens are to be used either for potentially sensitive applications, such as therapy or gamete generation, or for uses that were unknown at the time of consent. Data on participant attitudes in the field of genomics supports a reconsent approach when samples and data are being used for substantively different purposes than originally proposed [133]. There is also debate about whether pediatric participants should be recontacted to seek consent once they reach the age of majority [134]. All of this must be weighed against the
administrative burdens that reconsent provisions may have, including the potential of such provisions to slow down or otherwise impede needed research (especially if particular therapeutic benefit is found in a cell line but the initial participant cannot be found to get the needed permission).

With pluripotent stem cell research, the need for recontact does not end with future research permission. It may also be appropriate to recontact a participant for a variety of different reasons: disclosing incidental or individualized research findings, soliciting updated health information from the participant (e.g., to verify a specimen’s safety for human transplantation) [110–112], recruitment for future studies, describing changes in the scope of the current research study, and ensuring continued consent for use of a specimen obtained from a pediatric participant.

Reassurance
A consent form can be used as a tool to assure participants of the ethical use and governance of their specimen. However, providing detailed information about what will not be done with a participant’s specimen can be problematic. It may be sensible to reassure a participant that certain uses are prohibited, specifically reproductive cloning and germ-line introduction. Going beyond this, however, may facilitate misconceptions and may hinder research in unintended ways.

**Our Approach to Consent for iPSC Research**

The consent form template that we developed is included as supplemental online data, and the general structure is described below in Tables 3 and 4. The following sections provide the rationale for the content and structure decisions that we made, including provisions for future research uses, sustained interactions with participants, prospects for benefits and payments, and withdrawal from research, as well as logistical and governance issues. This analysis, at the very least, represents the necessary categories that must be addressed in formulating an iPSC consent process. We recognize that institutions and states will have specific regulations and requirements that may conflict with our recommendations; the consent form will have to be tailored appropriately in each case.

Our recommendations rely on several assumptions about the direction of the fields of iPSC research and regenerative medicine. First, as noted earlier, there is an intense focus on prospective planning to rapidly translate these cell lines to clinical use, both for drug development platforms and for cell transplantation and tissue engineering (Table 1). Second, we believe that the iPSC research enterprise will benefit from longitudinal relationships between specimen donors and researchers, supported by the fact that banking is most useful and ethical when ongoing commitments predicated on a thorough up-front consent can be honored.

**Future Research Uses**

The approach we take in addressing the scope of iPSC research (Table 3) is geared toward an “omnibus” consent model in which the research purposes are open-ended and include broad future uses and applications, within limits. Our framework draws a line past which prospective consent cannot go at this time. This line is structured such that cases of future problematic research will be relatively rare but would require recontact for reconsent. Our template assumes that some future research uses will require affirmative reconsent, including (a) reproductive research and (b) particularly sensitive areas of research that have not yet been anticipated. An inability to get reconsent from participants would proscribe using their iPSC lines for these research uses. We also followed the lead of some existing consent models and encourage participants to provide updated contact information so that ongoing contact will be possible.

We assume that including therapeutic goals—specifically, regenerative medicine and cell transplantation—is ethically permissible within the broad scope of a prospective consent, given that translation is one of the fundamental goals of iPSC research. This is a potentially controversial stance, and it is not the only approach currently being taken. Others have suggested that, like reproductive research, introducing derivatives of iPSCs made from a participant’s samples directly into other patients has special status and thus should require specific reconsent in the future [67], and this approach was reflected in some of the existing consent forms that we reviewed. However, there is precedent for consent approaches that clearly describe the prospects for therapeutic applications of iPSCs so that subjects can decide whether they are comfortable accepting those possibilities.

**Sustained Interaction**

Our consent template reflects an intention for sustained interactions [84] with participants in select circumstances about the ongoing uses of their coded specimens. This will be important for both the original and secondary researchers, as the need for recontact increases with time and scientific advancement. Provisions that support a sustained interaction include a description of plans for recontact in multiple but limited circumstances, the coding and linkability of specimens, and plans for engaging pediatric participants once they reach the age of majority (Table 4).

Our review of existing consent forms revealed significant variation in plans for recontact, including plans to de-link, which may be problematic as previously discussed. Our approach is consistent with the scope of permissible future research discussed above, permitting cell lines to be used for both research and translational applications and allowing researchers to meet obligations to provide participants with feedback over time.

Sustained interactions are also relevant in the case of pediatric subjects, who eventually will become adults over the course of the research and will have autonomous interests in the use of their specimens at that time. Accordingly, we have incorporated plans for pediatric reconsent that allow children to be informed upon reaching adulthood of the ability to review the previous consent. We endorse pediatric repository models that notify children when they reach the age of majority that their specimen is being used in research and that any decision making regarding the future use of those specimens, withdrawal, etc. now falls to them; we realize, however, that there might be difficulties in locating and tracking these participants [134, 135].

We acknowledge that our recommendations for sustained interactions with participants lead to additional logistical steps, such as developing web interfaces and newsletters that inform participants of the types of research being done, the results, the implications, and the goals. This increased communication will inform participants of the research to which they are integrally
Abbreviation: iPSC, induced pluripotent stem cell.

...participant has not consented to these projects, and other samples must be obtained and used. 

...conceivably be covered under the original consent (i.e., as Foundational research and applications in the table), as long as the distinction is explained.

...found some problematic language that hints at a possibility of questions is far from settled and because individualized findings may not be valid after cell lines have mutated or been reprogrammed, we have not taken a position on what (if any) obligations exist to return findings in this context, only facilitating the diversity of approaches to disclosure.

Prospects for Benefits and Payments

Given that the promise of iPSC research carries with it heightened expectations about the translational benefits and a risk of therapeutic misconception because of the emphasis on “personalized treatments” in public discourse, as well as questions about future commercial potential and patenting, our consent form template attempts to temper participants’ expectations [105, 136–139]. (Therapeutic misconception is the mistaken belief that research is both designed for and likely to help research subjects personally. It is common among research participants. The misconception is concerning because it might lead participants to believe that research on their samples will be prioritized for their particular conditions and that treatments will be developed for them [or their families] personally as part of the research study in which they are participating.) The template explicitly informs participants that direct benefits to them are unlikely and that they will not benefit financially from any commercial product developments based on their specimens (Table 4). We also included language underscoring that participants will not be able to retrieve cell lines developed from their specimens for personal use, nor can they choose who can receive future developments based on their specimens (Table 4). We also identified variation in the description of plans for the management and disclosure of incidental research findings. Because the ethical discourse on these topics that direct benefits to them are unlikely and that they are participating. For example, allowing withdrawal of just the identifying information associated with a specimen can help preserve some of the scientific utility of that specimen. Some existing consent forms describe more extensive opportunities for the withdrawal of original or derivative materials, which can be justified as long as these promises can reasonably be tracked and honored.

Withdrawal from Research Participation

Our approach permits various forms of withdrawal at different stages of the research process (Table 4). Although the logistics of a staged approach are more complicated and require tracking mechanisms, it yields a net benefit for both research participants and the research process. For example, allowing withdrawal of just the identifying information associated with a specimen can help preserve some of the scientific utility of that specimen. Some existing consent forms describe more extensive opportunities for the withdrawal of original or derivative materials, which can be justified as long as these promises can reasonably be tracked and honored.

Logistics and Governance

Our group struggled with how many separate decisions to grant participants within the consent form. Should participants be asked to agree to ongoing research as proposed, given the opportunity to agree to ongoing contact in certain cases, and/or given explicit choices (via check boxes) about the various aspects of the research? We attempted to carve out the simplest and broadest system we could justify, keeping in mind that check boxes, although attractive, could be logistically difficult to track and also may register participants’ reflexive reactions rather than their well-considered values. Any provisions allowing for explicit participant decisions that include check boxes necessitate a tracking mechanism. Although burdensome, it may be beneficial for iPSC banks to develop a standardized system to track and interpret these
Table 4. Other consent recommendations

| Domain                                      | Content                                                                 |
|---------------------------------------------|-------------------------------------------------------------------------|
| Sustained interactions with participants    |                                                                         |
| Managing individually identifiable information with “traceable” coded samples | 1. Confidentiality-related risks                                       |
|                                             |   • Unique nature of genome and its identifiability                     |
|                                             |   • Emotional and psychological risks related to third party disclosure |
|                                             |   • Stigmatization and other group risks                               |
|                                             | 2. Confidentiality protections                                         |
|                                             |   • Coding and other security measures                                  |
|                                             |   • Limits/exceptions                                                   |
|                                             |   • Legal protections (e.g., GINA, Certificates of Confidentiality)     |
|                                             | 3. Sharing of iPSC lines with other investigators                       |
|                                             |   • Whether/how other investigators using the specimens will be able to recontact participants |
|                                             | 1. Circumstances in which recontact may occur (including provisions for soliciting participants’ preferences for ongoing contact, if applicable) |
|                                             |   • To obtain reconsent for certain research uses (see Table 3)         |
|                                             |   • To solicit an additional specimen and/or updated health information from the donor that, e.g., might affect the use of the specimens in clinical interventions |
|                                             |   • To return individual research results (if applicable)               |
| Recontact                                    | 2. Necessity of updated contact information from participants           |
|                                             | 1. (When parents are authorizing participation of their children.) At the age of 18, enrolled children will be able to make any ongoing decisions regarding the research themselves\textsuperscript{a} |
| Pediatric samples                           |                                                                         |
| Benefits and interests associated with iPSCs |                                                                         |
| Future medical and societal benefit          | 1. Direct benefit                                                       |
|                                             |   • No direct benefit for subject or family                             |
|                                             |   • Participants should not expect individualized results               |
|                                             |   • iPSC banks set up for research are not equivalent to cord blood banks for personal use; cell lines cannot be retrieved by participants from researchers for personal use |
|                                             | 2. Societal benefit                                                     |
|                                             |   • Long-term research goals to better understand various diseases and develop better treatments in the future |
| Financial stake                             | 1. Commercial profit and financial stake in discoveries                |
|                                             |   • Participants will not receive any profits or royalties resulting from the use of their specimen, although it is possible that a company will profit from results of studies using the specimens |
|                                             |   • Disclose any known commercial interests of the researchers, institutions, sponsoring companies, and referring physicians (if appropriate) |
| Governance of specimens and withdrawal      | 1. Participants may request that all original materials not involved in an active research project be destroyed (skin, blood, hair, unmodified cell lines) |
| Scope and limitations of ability to withdraw from participation in iPSC research |   • Original material being used in a current project (as a control, for example) may remain in use |
|                                             | 2. Once specimens have been modified as part of a research project (gene or chemical modification, specifically including iPSC reprogramming) or banking initiative, these derivatives cannot be withdrawn or destroyed |
|                                             |   • Broadly distributed and cannot be retrieved                         |
|                                             |   • Participants may request that codes be removed so that specimens, cell lines, and information cannot be linked to identifying information |
|                                             |   • Cell lines can still be distributed in a deidentified fashion      |
|                                             |   • Participants can no longer be recontacted by the researchers        |
|                                             |   • Institutions who are known to have received any materials through distribution will also be asked to remove codes |
|                                             | 4. Participants may withdraw their consent to be recontacted while allowing their coded specimens and information to be used in research (if applicable) |
| Storage and banking\textsuperscript{b}      | 1. Purpose of a repository/“stem cell bank”                             |
|                                             | 2. Possibility that samples, iPSC lines, and associated data may be placed in repository |
|                                             | 3. Information about repository’s governance and review policies regarding, e.g., how it is decided who may receive cells and data |
|                                             | 4. Timeline for banking (e.g., “indefinitely”) and for using contributed specimen to create and distribute iPSCs, if known |
| Limitations on participant choices           | 1. Participants have no control over who may benefit from or receives treatment derived from research on their specimens |
|                                             | 2. Participants cannot delineate limits on which diseases can be studied with their specimen |
|                                             | 3. Specimens may be used to study diseases that are outside of the scope of the original study |

\textsuperscript{a}Some protocols will notify pediatric participants when they reach 18 years of age regarding their ongoing research participation, allowing them to opt-out if they wish. All future recontact would be directed at the now-adult participant. Such provisions could also be described to parents in the consent form in more detail.

\textsuperscript{b}Appropriate tracking mechanisms for participants’ contact information, consent preferences, etc., are an important component of these provisions. Abbreviation: iPSC, induced pluripotent stem cell.

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understanding of the complexities of iPSC research, we recommend that supplementary educational materials describing iPSCs be distributed to participants along with the consent form. The manual developed by the International Society for Stem Cell Research [141] is a useful model.

CONCLUSION

Approaches to informed consent for iPSC research thus far have been variable, often in ways that could create confusion for participants and hinder research collaborations and future uses. This variation goes beyond what we might expect based on differences in study design and institutional policies; it seems to reflect fundamental disagreements about the appropriate scope of the consent process at this time, the manner in which future plans should be described, and the necessity of ongoing contact with participants. This lack of consensus is problematic in a field with accelerating momentum that will rely on the sharing of cell lines.

Our proposal reflects the broadest possible approach to consent that we believe can be ethically justified, incorporating all of the currently foreseeable ways in which iPSC research will play out. Although our approach does risk excluding some participants who are not comfortable with the stated long-term goals, we accept this as the trade-off for making a model that is practically useful to researchers as well as to a majority of research participants.

On its surface, our reappraisal of informed consent for iPSC research may seem to have limited consequences. It is important, however, as a first step in the larger process of improving dialogue between the research enterprise and the public. Although empirical data are limited, there is some evidence that the public has visceral discomfort, both legitimate and unfounded, with certain forms of research. As Aalto-Setälä et al. warn, “if the perception that iPSC research poses no ethical concerns is not corrected, there could be a backlash against iPSCs later” [67].

The importance of informed consent extends beyond the physical, economic, and psychosocial risks associated with iPSC research [72]. Research participants have autonomy-based interests in the longitudinal uses of their samples, especially when they might object to some uses on moral grounds. Thoughtful informed consent, accordingly, plays a role that is critical to the success of research with iPSCs: assuring the public that researchers will “honor both the letter and spirit of the agreements between researchers and subjects” [72]. We believe that the proposal described herein represents a significant step toward these goals.

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AUTHOR CONTRIBUTIONS
J.L.: conception and design, data analysis and interpretation, manuscript writing, final approval of manuscript; S.L.: conception and design, final approval of manuscript, liaison to investigators during consent form drafting process; M.R.: conception and design, administrative support, final approval of manuscript; S.C.H.: conception and design, financial support, administrative support, provision of study material, manuscript writing, final approval of manuscript.

DISCLOSURE OF POTENTIAL CONFLICTS OF INTEREST
The authors indicate no potential conflicts of interest.

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