The ethical anatomy of payment for research participants

Joanna Różyńska

Accepted: 4 May 2022 / Published online: 24 May 2022 © The Author(s) 2022

Abstract
In contrast to most publications on the ethics of paying research subjects, which start by identifying and analyzing major ethical concerns raised by the practice (in particular, risks of undue inducement and exploitation) and end with a set of—more or less well-justified—ethical recommendations for using payment schemes immune to these problems, this paper offers a systematic, principle-based ethical analysis of the practice. It argues that researchers have a prima facie moral obligation to offer payment to research subjects, which stems from the principle of social beneficence. This principle constitutes an ethical “spine” of the practice. Other ethical principles of research ethics (respect for autonomy, individual beneficence, and justice/fairness) make up an ethical “skeleton” of morally sound payment schemes by providing additional moral reasons for offering participants (1) recompense for reasonable expenses; and (2a) remuneration conceptualized as a reward for their valuable contribution, provided (i) it meets standards of equality, adequacy and non-exploitation, and (ii) it is not overly attractive (i.e., it does not constitute undue inducement for participation or retention, and does not encourage deceptive behaviors); or (2b) remuneration conceptualized as a market-driven price, provided (i) it is necessary and designed to help the study achieve its social and scientific goals, (ii) it does not reinforce wider social injustices and inequalities; (iii) it meets the requirement of non-exploitation; and (iv) it is not overly attractive. The principle of justice provides a strong ethical reason for not offering recompenses for lost wages (or loss of other reasonably expected profits).

Keywords Research ethics · Payment for research participants · Social beneficence · Autonomy · Justice/fairness · Undue inducement · Exploitation

What are the ethical principles or values which constitute an ethical rationale for paying research subjects? Do those create a moral obligation to pay individuals for participation in biomedical research, or rather a mere justification for its acceptability? What other ethical principles and values shape the payment practice, and how? Surprisingly, these questions have been rarely the subject of in-depth discussions in the literature. Instead of exploring the ethical foundations of payments systematically, scholars and public-policy makers rather focus on payment-related ethical concerns, in particular, risks of undue inducement and exploitation, and—from this perspective—recommend or discourage certain forms, schedules and timings of payment commonly used in research practice (e.g., Macklin 1981; Dickert and Grady 1999; Grady 2001, 2005; Gelinas et al. 2018; Resnik 2015, 2019; Largent and Lynch 2017a, 2017b). As a consequence—while there is a growing consensus that an ethically sound payment scheme should avoid both excessive payment and underpayment, and it should include, at least, reimbursement of reasonable expenses and compensation for some contributions made by research subjects—there is no generally accepted view on whether a payment to research subjects (as such or of a certain kind) is a moral obligation (Council for International Organizations of Medical Sciences 2016), merely an “acceptable practice” (Food and Drug Administration 2018), “ethically discretionary” activity (Persad et al. 2019, p. 319), or just a “necessary evil”. Equally, there is no common view on what constitutes an ethical source of this purported obligation or acceptability of payment (as such or of a certain kind), and which ethical reasons lie behind different payment categories and schemes.

This paper aims at clarifying these issues. It presumes that any discussion on ethically sound payment practice should be preceded by a clear statement of ethical reasons...
for paying research participants, their deontic nature and mutual relations. Without full understanding of the ethical anatomy of payment, it is impossible to determine what we owe, if anything, to research subjects—what for, and how much research participants should be paid.

Preliminary terminological remarks

Paying research subjects for their participation in biomedical studies is an increasingly common practice across different types of research involving healthy volunteers and patients (Grady et al. 2005; Largent and Lynch 2017a). Nevertheless, the payment continues to raise numerous conceptual, ethical and practical controversies among bioethicists, investigators, research ethics committees/institutional review boards (RECs/IRBs), and other members of the research community. Although prominent international guidelines and national regulations call attention to the crucial moral issues that payment raises (in particular, the risk of undue influence), they offer little substantive guidance on how to pay research subjects in an ethical way, and if they do so, they often provide contradictory advice. For instance, the World Medical Association Declaration of Helsinki (2013) does not address payment for research participation directly. It only mentions that information on “incentives for subjects” must be contained in the study protocol (par. 22). Also, the U.S. “Common rule” (Department of Health and Human Services 2018) and the European Union “Clinical Trial Regulation” (2014) offer very limited guidance on payment for participants. All these regulatory lacunas and contradictions are reflected in considerable variation in local payment policies and practices around the world (Dickert et al. 2002; Grady et al. 2005; Fry et al. 2005; Ndebele et al. 2008; Pasqualetti et al. 2010; Roche et al. 2013; Largent and Lynch 2017a).

One of the factors which adds to this confusion is the diversity and ambiguity of terminology used in the literature and guidelines on research payment. For example, the International ethical guidelines for health-related research involving humans of the Council for International Organizations of Medical Sciences (CIOMS) make a distinction between “reimbursement” for reasonable direct costs incurred by research subjects and “compensation” for the time spent and other inconveniences resulting from study participation (2016, Guideline 13 and Commentary). The Guideline for good clinical practice issued by the International Conference on Harmonisation of Technical Requirements for Registration of Pharmaceuticals for Human Use speaks of “payments and compensation” (2016, par. 3.1.2., 3.1.8., 3.1.9.) with no further explanation. The U.S. Food and Drug Administration Guidance for institutional review board and clinical investigators: payment and reimbursement to research subjects (2018) makes a distinction between “reimbursement” for direct costs and other “payment” for participation, which it considered to be more ethically challenging as in some cases it may constitute an undue influence to prospective subjects. The Council of Europe’s Additional Protocol to the Convention on human rights and biomedicine, concerning biomedical research mentions „payments and rewards” without any additional differentiations, in the Appendix containing a template of an information sheet for the ethics committee (2005a). The term is “unpacked” in the Explanatory Report to Article 12 of the Additional Protocol dealing with undue inducement which refers to “compensation” for burdens and inconveniences and “reimbursement” for expenses and financial losses (Council of Europe 2005b, sec. 64). Australian guidelines on Payment in participation in research: information for researchers, HRECs and other ethics review bodies issued by the National Health and Medical Research Council (2019) divide “payment” for participants in four analytical categories: “reimbursement” for any research-related expenses; “compensation” for any documented financial losses resulting from participation in research, including loss of wages, or from an injury suffered as a direct consequence of participation; “remuneration” paid to participants in recognition of their service for the time spent and other inconveniences resulting from participation; and “incentive or inducement” provided to individuals simply to encourage their enrolment or continuation in research. In the United Kingdom, the NHS Health Research Authority guidance (2014) follows terminology developed by the Nuffield Council on Bioethics. The Council, in its report Human bodies: donation for medicine and research (2011) distinguishes three forms of payment: (i) “recompense” offered in recognition of losses incurred which may take a form of “reimbursement” of direct financial expenses, lost earnings, or “compensation” for non-financial losses, such as time, inconvenience and discomfort; (ii) “reward” defined as a “material advantaged gained by a person …that goes beyond ‘decompensating’ the person for losses they incurred”; “reward” becomes “remuneration” when calculated as a wage or equivalent”; and (iii) “purchase”—money given in exchange of a “thing” (e.g., biological material for research) (Nuffield Council on Bioethics 2011, p. 2).

In order to bring an end to this terminological (and conceptual) chaos, in this analysis, “payment” is used as an overarching term that encompasses all forms of financial and in-kind support provided to research participants.1 It covers two sub-categories, which will be referred to as “recompense” and “remuneration”. These categories are distinguished by a different impact each of them has on the participant’s economic position as evaluated ex post.

---

1 This is an improved version of the payment typology presented in Różyńska (2021).
The term “recompense” stands for any payment that entails no net benefit to recipients. Recompense amends to research participants for financial and non-financial losses or injuries resulting from their participation in research. Thus, this type of payment does not constitute a gain or profit, but merely covers—understood in broad terms—costs of the participation. Recompense may include three sub-categories of payments: (i) reimbursement of direct financial expenses incurred as a result of participation in research (e.g. costs of travel, accommodation, meals, childcare); (ii) compensation for indirect financial expenses, i.e., losses, that arise from participation in research (e.g. loss of wage in result of taking unpaid leave from work); and (iii) compensation for financial and non-financial losses resulting from injuries suffered as a direct consequence of participation in research.

“Remuneration” refers to any payment provided to individuals for their service as research participants, which exceeds expenses, losses or injuries experienced by participants as a result of their participation, and brings net benefit (gain, profit) for recipients. The remuneration for research participation (same as for “regular work”) may be understood either as a reward given in recognition of and as an appreciation for participants’ valuable contribution; or as a price—money given in direct, market-driven exchange for a service provided by participants (cf. Moriarty 2020; Różyńska 2021). If the remuneration is understood as a reward, the payment should be adequate to the desert, i.e., to the value of participants’ contribution. The latter value necessarily depends on various factors, including (i) time allotted to research; (ii) efforts or types of services rendered (e.g., performing psychological or physical tests, taking drugs or using device as instructed, conducting self-monitoring or gathering other research-relevant information); (iii) discomfort, burdens or inconveniences associated with participation (e.g., stress, pain, suffering, but also burdens related to sticking to a dietary regime or inconveniences caused by the expected lifestyle changes); (iv) the level of risk involved in research, and (v) special/unique value of the input of the specific subject (e.g. due to rarity of a disease suffered by the person or her genetic make-up). If remuneration is considered as a price for participation, the amount of payment is determined by market forces—supply and demand, and it does not have to be proportionate to the value of the subjects’ contribution. In literature and guidelines, the latter way of thinking about payment if often concealed under the concept of “incentive” or “inducement”, although—as it will be explained later—these labels are far from being adequate and should be abandoned.

The above typology of payments for research participation is preferable over other schemes proposed in the literature and regulatory documents at least for three reasons. First, it is built upon one clearly defined, objective and disjoint divisional criterion, thereby avoiding a mistake of mixing entirely different criteria in one scheme. Such a mistake affects all typologies of research payments which, alongside recompenses and remunerations, distinguish “incentives” or “inducements” payments. This mistake originates from the fact that the former two categories of payments are defined by the payment impact on the subject baseline economic position, the latter is based on the researcher’s intention. Secondly, by rejecting the term “compensation”, the proposed typology avoids conceptual and normative confusion stemming from ambiguity of this notion in legal terminology, where it stands for both money received in return for services rendered, especially salaries or wages, and for payment of damages for loss or injury (Różyńska 2021). The ambiguity undermines the conceptual and normative value of highly popular payment scheme advocated by Gelinas, Largent, Lynch, and collaborators (Gelinas et al. 2018; Persad et al. 2019; Lynch et al. 2021; Bierer et al. 2021). Following Christine Grady’s terminology (2005), the authors separate “reimbursement”, “compensation”, and “incentive”. Trapped by the ambiguous language of “compensation”, they frame remuneration as recompense for losses, arguing that “participants’ time, as well as their assumption of research-related burden and inconvenience … are critical contributions experienced as losses by participants; they are giving up or accepting unfavorable as a result of participation. Thus, the same justifications applicable to reimbursement applied here” (Lynch et al. 2021, p. 16). At the same time, applying the logic of remuneration, they reject compensating participants based on their opportunity costs, noting that “rather, research compensation is an acknowledgement of participants’ contributions to research, which leads to the conclusion that there generally should be equal pay for equal work” (Lynch et al. 2021, p. 16).

Finally, the advocated typology allows for separating payments form small gifts given at the conclusion of a study (such as chocolates, T-shirts, cups, pens, cinema-tickets) that have minimal market value, serve only as token of appreciation, and have likely zero impact on recruitment (Grady 2005).

Having clarified basic terminological and conceptual issues, we can move on and explore the ethical anatomy of payment for research participants. It would be useful, however, to indicate briefly—from the outset—the scope of the forthcoming analyses.

The paper deals with the widespread practice of paying research participants in exchange for their valuable service without determining whether the service should be treated as an unskilled labor, “regular” work, body renting, or a unique sui generis endeavor (cf. Lemmens and Elliott 1999, 2001; King Reame 2001; Anderson and Weijer 2002; Elliott 2008; Abadie 2015; Phillips 2011b; Lynch 2014; Różyńska 2018). For the sake of the analysis to come, it is assumed that paid participation in biomedical research is a form of
paid bodily services, and it should be “no more worrisome to commodify a person’s labor [bodily service—JR] as a research subject than to commodify a person’s labor in other contexts, which happens all the time” (Lynch 2014, p. 159). Therefore, commodification concerns against research payment, raised by some commentators (Macklin 1989; Chambers 2001; Abadie 2010, 2015; Cooper and Waldby 2014; Walker and Fisher 2019) will not be explored here.

Since the paper focuses on the most fundamental ethical basis of payment practice, it is limited to monetary offers (paid in cash, cheques or pre-paid credit cards) as a paradigmatic case of payment, leaving aside all forms of in-kind support. This paper also leaves aside payments to participants unable to give consent, because those raise additional, substantive issues, as well as recompenses for research-related injuries as they would require in-depth legal analysis. The presented analysis applies to research which RECs/IRBs would consider scientifically and ethically sound thus presenting arguments that payment is offered to overshadow or mask some ethical deficiencies of a study project, especially those regarding its risk–benefit profile or criteria of subjects selection. In other words, it considers “research which ethics committee would allow to proceed, were the subject not paid” (Wilkinson and Moore 1997, p. 375).

Ethical rationale for paying research participants

The main “pragmatic” reason (Largent and Lynch 2017a, p. 77 fn. 22) for offering payment to research participants is to boost recruitment and retention rates. This “efficiency”-driven (Phillips 2011a) rationale is widely acknowledged. It is, however, rarely recognized as having not only practical, but also an ethical facet. Since progress in biomedical science and healthcare is not ethically discretionary, this section argues that neither efficient enrolment nor payment for research subjects, being conductive to the latter, are value-neutral practices. On the contrary, they are ethically grounded in and governed by the principle of social beneficence that calls for maximization of a common good, i.e. good which has two characteristics: it is non-exhaustible (one person’s use does not diminish another’s use) and is beneficial for all or almost all members of a society.²

Biomedical research is social practice aimed at generating such a common good—generalizable scientific knowledge that may contribute to the improvement of healthcare interventions and public health measures, thus leading to the advancement of human health (cf. Schaefer et al. 2009; Rhodes 2010; London et al. 2010). Health is highly valued by all individuals and societies primarily “because of what it enables us to do” (Duncan 2010, p. 321). A minimum threshold of physical and mental capacities is necessary (though not sufficient) for an individual to be able to pursue her particular life goals, plans, and projects that express her vision of a good life. Since all liberal and democratic societies share a fundamental moral (and political) commitment to protecting and respecting each person’s right to lead her life in accordance with her personal views of what is valuable in human life, they also share a moral obligation to encourage forms of social collaboration useful in fulfilling basic health needs of their members (London 2003, 2006; Rawls 1971; Nussbaum 2013). This includes the practice of healthcare as well as biomedical research, because the capacity of a society to satisfy its members’ basic health interests is conditional upon progress in biomedical sciences, which in turn is crucially dependent on various types of research, including human research. Thus, societies have a prima facie moral obligation to promote the conduct of biomedical research, including research involving human subjects.

Human biomedical research is a complex, collaborative social “enterprise”, involving various institutional and individual stakeholders with various, often conflicting, interests (London et al. 2010; Różyńska 2015). Therefore, in order to fulfill their obligation to promote biomedical research, societies should develop, implement and support normative and institutional mechanisms aimed ensuring the existence, stability and effectiveness of research practice in attaining the common good (London et al. 2010; Resnik 2011; Różyńska 2015). These mechanisms should enhance all stakeholders’ trust and willingness to support and invest in the research enterprise, including prospective research participants without whom biomedical research would not be able to achieve its goals.

Socially valuable human biomedical research is critically dependent on successful enrolment and retention of a sufficient number of appropriate participants, and on their willingness to comply with study procedures and conditions. Failure to recruit or retain participants may lead to invalid or inconclusive research data, it may result in

---

Footnote 2 (continued)

members of the community… interests in being able to develop their intellectual and affective capacities in order to pursue activities that they find meaningful, and to engage in meaningful relationships with others” (2003, p. 21).
Regarding financial costs and benefits linked to research parity requirements, seems to be the most important, concerns and clinical character, such as access to trials and eligibility and retention rates. Although barriers of structural and clinical character, such as access to trials and eligibility requirements, seems to be the most important, concerns regarding financial costs and benefits linked to research participation also play a significant role in prospective subjects’ decision-making (Friedman et al. 2015; Hamel et al. 2016; American Cancer Society Cancer Action Network 2018; Nipp et al. 2019a, Unger et al. 2013, 2019). Recent statistics published by the Center for Information and Study on Clinical Research Participation show that information about potential costs and their reimbursement as well as information about compensation for time off from work are among the most important factors influencing a decision to participate in research for—respectively—58% and 40% potential subjects (2019). And the prospect of receiving monetary compensation is one of three top reasons (34% mentioning) impacting a decision to enroll into a study (Center for Information and Study on Clinical Research Participation 2019). These data are consistent with results of numerous empirical studies on subjects’ motivations for volunteering, conducted among patients and healthy volunteers. They all indicate that although the payment is not the only reason why people agree to participate in biomedical research, it is definitely one of the top motivates for enrollment, especially among healthy volunteers (Tishler and Bartholomae 2002; Almeida et al. 2007; Abadie 2010; McCann et al. 2010; Stunkel and Grady 2011; Grady et al. 2017; Fisher et al. 2018; Manton et al. 2019).

Thus, although it is rightly stressed in the literature that the exact impact of paying for participation on the rates of research enrollment and retention remains still under-investigated, it is reasonable to assume that payment can make a positive difference in this respect (Dunn and Gordon 2005; Watson and Torgerson 2006; Caldwell et al. 2010; Probstfield and Frye 2011; Treweek et al. 2013; Nipp et al. 2019b; Parkinson et al. 2019). Money may remove participation barriers for those individuals who are unable or unwilling to cover direct costs associated with research, such as costs of traveling, lodging or hiring a babysitter. The payment may attract people who would be otherwise discouraged from the participation by the necessity of taking unpaid leave from work and resulting loss of wages. It may help to make a positive enrollment decision for those, who feel forced by their low social-economic status (SES) to dedicate time and efforts for searching for a job or earning their living, instead of altruistically contributing to the development of science. Additionally, an offer of payment could convince to participation persons who believe that their private or professional time is particularly valuable (for example due to their unique responsibilities, competences, or skills), and, therefore, they should not allot it to alternative causes without an adequate remuneration. In all these cases, payment may enhance recruitment, provided it is designed in a way that targets the underlying barriers, needs, or expectations.

Insofar as these factual assumptions about inducing potential of money are true—and we concede them for the sake of the argument—the principle of social beneficence provides fundamental ethical rationale for paying research subjects. Offering payment to participants is an ethically right and prima facie obligatory practice because it increases research recruitment and retention, thereby contributing to the common good produced by the research enterprise. From a broader perspective, “it is useful in fulfilling society’s obligation to meet the essential [health-related] needs of its members” (Ackerman 1989, p. 1). The strength of the obligation to offer payment for participation grows in relation to studies which are urgently needed, e.g., to address acute public-health emergencies (such as a dire pandemic), or when there is strong evidence that without payment recruitment, retention and completion of socially valuable studies would be doomed or severely compromised. The strength of the obligation weakens when—due to study-specific features—payment offers are not needed to secure an adequate number of participants, i.e., where it is reasonable to assume that people will be willing to join the study for non-economic and social reasons, e.g., the prospect of direct or ancillary medical benefits, the wish to make a contribution to medical progress or the health of others, scientific curiosity, interest in the goals of the study, the prospect of making friends or having new experiences (Stunkel and Grady 2011; Grady et al. 2017; Fisher et al. 2018; Manton et al. 2019).
Additionally, the obligation to pay research participants may be overridden by justified research budget constraints. If investigators have no money to pay for participation in a socially valuable research project, it is better to allow them to proceed without payment, than obliging them to pay, thereby forcing them to resign from conducting the study (Gelinas et al. 2018).

To sum-up, the payment for research subjects is an incentive or inducement for participation grounded in the principle of social beneficence. This is *expressive verbis* acknowledged by the U.S. Food and Drug Administration (2018) guidance which argues that payment to research subjects is “a recruitment incentive” and by the European Union’s Clinical Trial Regulation (2014) which refers to any payment offered to trial participants as “incentives or financial inducements” (art. 31.1(d); art. 32.1(d), 33(d)). Thus, payment is not, as some scholars and guidelines suggest, a demand of justice or fairness (Gelinas et al. 2018, 2020; Persad et al. 2019; Lynch et al. 2021) or requirement of non-maleficence and beneficence for an individual subject (Bierer et al. 2021). These principles, supplemented by the principle of respect for autonomy and other considerations regarding non-exploitation, are nevertheless very important. While they do not provide an ethical rationale for offering payment for research participations by themselves, they set contours for an ethically sound payment practice.

**Ethical contours of an ethically sound payment practice**

Offering payment for research participants is an incentive *prima facie* required by the principle of the social beneficence. This does not imply, however, that it is ethical to set payment at whatever level necessary and sufficient to attract an adequate number of proper subjects in a timely fashion. On the contrary, no matter how payment is important and effective as an incentive for participation, not every amount, method and timing of payment is acceptable. The reason being the fact that the consequentialistic principle of social beneficence does not exhaust the reinig normative framework for human biomedical research. The ethics of biomedical research is built upon a matrix of principles and values which strives to find an adequate balance between the imperative to advance interests of science and society (“research imperative”) and obligations of all societies to protect other important interests of their members, especially interests of research participants and/or involved communities. These latter obligations may be viewed as constituting a general “imperative of non-exploitation” that sets boundaries for the practice of human biomedical research, in general, and for the practice of paying participant, in specific.

**Autonomy and payment**

One of the core values behind the imperative of non-exploitation in research is the value of human autonomy. The principle of respect for autonomy is a cornerstone of modern research ethics (National Commission for the Protection of Human Subjects of Biomedical and Behavioral Research 1979; Faden and Beauchamp 1986; Emanuel et al. 2000; Beauchamp and Childress 2001). It is also a bedrock of a fundamental ethical requirement for research, namely informed consent that serves to “ensure not only that individuals control whether or not they enroll in clinical studies”, but also that “they participate only when doing so is consistent with their values and interests” (Emanuel et al. 2000, p. 2706).

The principle of respect for autonomy has a high relevance for the practice of payment for research subjects as one of the most commonly expressed concerns is that payment can be coercive or constitute an undue inducement (undue influence), thus compromising the validity of informed consent (e.g., Macklin 1981; Faden and Beauchamp 1986; McNeill 1997; Dickert and Grady 1999, 2008; Largent et al. 2013; Gelinas et al. 2018, 2020). It is claimed that very high or (for other reasons) overly attractive payment may undermine the capacity of individuals to make autonomous decisions regarding study participation, by compromising their voluntariness, ability to adequate understand and assess research risks and benefits, or by “forcing” them to make choices against their “better judgment or deeply held beliefs” (Council for International Organizations of Medical Sciences 2016, p. 54).

Although there is still a substantial disagreement in the literature about whether an offer of payment may be perceived as coercion, what exactly constitutes undue inducement, and whether money can distort or compromise autonomous decision-making (Macklin 1981, 1989; Wilkinson and Moore 1997; Grady 1999, 2001, 2005; Grant and Sugarman 2004; Emanuel 2004, 2005; Wertheimer and Miller 2008; Klitzman 2013; Largent et al. 2013; Resnik 2015, 2019; Belfrage 2016; Largent and Lynch 2017a, 2017b; Millum and Garnett 2019), existing guidelines and regulatory documents disallow payment, which may unduly seduce people to consent for participation (Largent and Lynch 2017a). Thus, even though overly attractive payment offers could enhance timely recruitment by “alluring” a sufficient number of adequate individuals to join and stay in a study even against their better judgment, there is a regulatory consensus that such payment should not be offered.

The ethical unacceptability of such an overly attractive payment stems from the respect for autonomy. However, it also finds support in the principle of social beneficence. Common sense and empirical evidence, although still limited, suggest that attractive payments may have a negative
impact on scientific value of research, as it may encourage potential and/or actual participants to withhold or misrepresent information, which are critical for their recruitment eligibility or continuation in research (Bentley and Thacker 2004; Dresser 2013; Dickert 2013; Devine et al. 2013, 2015; Largent and Lynch 2017a; McManus and Fisher 2018; Lynch et al. 2019). Such a concealment, fabrication or falsification by participants create risks for participants, but also for research resources and the integrity of research data as it can bias the results and undermine the validity of a study (Lee et al. 2018). And—if it occurs frequently enough—it may jeopardize the whole research enterprise. Therefore, assuming—what stills needs to be explored empirically—a positive correlation between the prevalence of deceptive behaviors and the attractiveness of payment (which depends not only on the amount, but also on payment method and timing), the principle of social beneficience provides ethical reasons for employing payment strategies which do not involve overly attractive payment schemes.

The principle of respect for autonomy has fueled much of scholarly discussion on unethical nature of overly attractive payment. However, little attention has been paid to that principle in the context of no-payment and underpayment. Undoubtedly, the lack of resources to cover direct or indirect costs associated with the participation may constitute a barrier for individuals, who otherwise would be willing to take part in research (American Cancer Society Cancer Action Network 2018; Nipp et al. 2019a; Largent and Lynch 2017b; Gelinas et al. 2020; Bierer et al. 2021). The removal of these barriers by covering the relevant expenses is important for promoting potential subjects’ autonomy, because it enables individuals to exercise their free will to contribute to the development of science by serving as research subjects. Thus, the principle of respect for autonomy supports paying subjects remunerations, which make the participation in research a cost-free and revenue-neutral activity.

Finally, what is rarely observed, the principle of respect for autonomy provides a general support for public policies which allow remuneration of research subjects for their contributions—both in the form of reward and price—as it calls for respecting people’s right to decide freely in what practices and activates they what to engage in for the sake of earning their living. As Wilkinson and Moore note, “denying people the option of taking inducements reduces their freedom, since it removes an option that they prefer to the alternatives” (Wilkinson and Moore 1997, p. 377). Admittedly, biomedical research enterprise does not aim at broadening the scope of subjects’ freedom or autonomy. Nevertheless, it should not restrict people’s choices without a good reason. Participation in socially valuable biomedical research, whether or not one is paid for it, is neither morally wrong nor bad for the person concerned. On the contrary, it is considered at least as good as engaging in any other socially valuable and risky service or work. Therefore, there are no ethical grounds (either paternalistic or non-paternalistic) for depriving competent individuals an opportunity to serve as research subjects in exchange for money, provided that their decision to participate in a given study is autonomous (i.e., based on comprehensive and adequately understood information, and free from unduly controlling influences).

Moreover, some studies suggest that remuneration can in fact play a positive role in prospective subjects’ consent process by reducing the therapeutic misconception and highlighting research risks. Money might send a message to participants that they take risks and burdens for the sake of the benefit of science and society and “should be compensated for it, which would not occur if they were … expected to benefit from it.” (Glannon 2006, p. 252; also Dickert and Grady 1999; Grady 2001, 2005; Menikoff 2001; Largent and Lynch 2017a). Additionally, some studies suggest that payment can enhance autonomous decision-making by drawing prospective subjects’ attention to research risks and inconveniences (Cryder et al. 2010; Largent and Lynch 2017a; Fisher et al. 2019).

Justice, fairness and payment

Another ethical principle that has a direct relevance for the practice of paying research subjects is the principle of justice, both in its distributive and commutative dimensions. Many scholars worry that payment may be more attractive to individuals of lower SES, and thus offering payment for participation may result in unfair distribution of research benefits and burdens across the general population (e.g., Maclin 1981, 1989; Faden and Beauchamp 1986; Ackerman 1989; McNell 1997; Grady 2005; Dickert and Grady 2008; Gelines et al. 2018). The worse-off will shoulder a disproportionate share of burdens of research, while the benefits will accrue primarily to the better-off. A different, yet linked, concern relates to the risk of exploitation of research participants, especially those of low SES (Lemmens and Elliott 1999; Shamo and Resnik 2006; Elliott 2008; Elliott and Abadie 2008; Abadie 2010, 2015; Stones and McMillan 2010; Phillips 2011a; Resnik 2015, 2019; Largent and Lynch 2017b; Gelines et al. 2020; Bierer et al. 2021; MacKay and Walker 2021). Exploitation in research is about the unfair distribution of goods that arise from interaction between researchers and participants (Wertheimer 2008). “One party gets too little, while the other gets too much. Often, but not always, the unfair distribution arises because one party to the interaction is in a weak position, due to poverty, ignorance, or extreme urgency, which the other party can take advantage of, offering few benefits” (Emanuel 2004, p. 101). Economically disadvantaged individuals are in need of money and they tend to value a specific amount of payment more. For
many of them accepting an offer of unfairly low but still \emph{en bloc} beneficial payment for participation might be a reasonable choice. After all, even small payment is better than no payment at all, although it constitutes a (mutually beneficial and consensual) exploitation.

These concerns are highly important, but—as it is shown below—they do not exhaust the role of justice/fairness in shaping an ethically sound payment practice. Moreover, in order to fully understand their consequences for ethics of paying research subjects, it is essential to note two things.

Firstly, biomedical research is a social practice embedded in specific social reality shaped by its historical and cultural roots, reigning power- and economic relations, and normative fabrics. This social reality is marked by profound social and economic inequities (within concrete societies as well as at the level of international community), which should be mitigated, and ideally eliminated by broad social reforms. Research enterprise should not entrench or exacerbate these background social injustices. However, it is neither designed for nor capable of actively fighting them, especially against participants’ poverty, unemployment or lack of access to high-quality healthcare (Fisher 2019). Justice in research requires treating all research participants fairly and equitably, also when it comes to payment for their participation. But it does not require paying participants of low SES in order to alleviate or remove hardships of their position. Such a positive impact of research payment on subjects’ social or economic condition is laudable, and even desirable, but it is not a demand of justice.

Moreover, contrary to views of some commentators, the commutative justice per se does not require offering payments to research subjects. Largent, Emanuel and Lynch claim that “when goods and services are not indented as gifts, failure to pay for them is a problem: we call it theft” (2019, p. 1), thus suggesting that this is exactly what happens when participants are not fairly paid for their contribution to the common good. Despite its rhetorical attractiveness, this claim cannot be accepted as it rests on two mistakes. Firstly, it ignores the power of consent. Valid consent of a prospective subject for using her body for research purposes transforms theft into gift, lease, rent, work or other consensual relation with a researcher. Second, it forgets that it takes two willing parties to change provision of goods or services into transaction, i.e., exchange of goods and services in return for money. Thus, when a researcher is unable (e.g., due to budgetary constraints) or uninterested in offering payment for prospective participants (e.g., due to availability of sufficient number of unpaid subjects), or when a prospective participant is genuinely willing to contribute to the development of science without any remuneration and reimbursement, there is no ground for transaction. And there is nothing essentially unfair in allowing researchers and altruistically motivated participants to engage in scientifically and socially valuable biomedical research.

Payment is a recruitment incentive justified and \emph{prima facie} required by social beneficence, not by justice or fairness. However, \emph{when it is to be offered}, the offer should satisfy the requirement of distributive and commutative justice.

It is commonly accepted that the principle of justice requires distributing burdens and benefits of study participation in such a way that no segment of the population is unduly burdened by research or denied its potential or actual benefits. Recruitment criteria should reflect the scientific purpose of the study, not target populations which are considered “easy to recruit” “simply because of their easy availability, their compromised position, or their manipulability” (National Commission for the Protection of Human Subjects of Biomedical and Behavioral Research. 1979). Thus, researchers must neither exploit the vulnerable, in particular economically disadvantaged, nor exclude without good reason those who stand to benefit from study participation. They should strive to recruit an adequate cross-section of the population in order to spread research burdens and benefits fairly across the population. In other words, an adequate diversity of gender, race, ethnicity, age and SES in research should be sought (Geller et al. 2011; Kwiatkowski et al. 2013; Heller et al. 2014; Winter et al. 2018; cf. Dickert 2009; Bierer et al. 2021). It is worth noting that the last postulate finds an additional support in the principle of social beneficence as it enables the generalization of knowledge to be gained in research, thereby enhancing research scientific and social value (Wilkinson and Moore 1997; Grady 2005; Resnik 2015; Largent and Lynch 2017b). Thus, justice encourages payment schemes which have a potential of making distribution of research burdens fairer. And justice is against schemes that can deepen social inequalities, lead to unfair social distribution of research burdens and benefits or exploitation.

What should research subjects be paid for and in which amount to make payment consistent with demands of justice? Firstly, justice-related considerations provide an ethical justification for paying research subjects a recompense for direct costs related to research participation. By making participation in research a cost-free activity, recompense removes—at least some—economic barriers for participation, thus equalizing opportunities for all willing individuals to contribute to the development of science, no matter their SES. Moreover, as studies already referred to suggest (Unger et al. 2013, 2016, 2019; American Cancer Society Cancer Action Network 2018; Nipp et al. 2019b; Chino and Zafar 2019), reimbursement of direct expenses may improve access to potentially beneficial studies for patients, who otherwise would not be able to participate due to their low SES or disease-caused greater sensitivity to economic burdens associated with research. Thus, the reimbursement of
direct costs has a potential to enhance justice in research not only by reducing inequities in access to research for those who otherwise could not afford it, but also by contributing to the fair distribution of clinical benefits associated with participation.

Secondly, the principle of justice provides a strong ethical reason for disallowing recompenses for the loss of reasonably expected profits, especially lost wages. Such recompenses lead to differential payments between participants—individuals who receive higher wages get higher recompense; those who are lower paid by employees—lower recompense, and those who are unemployed—no recompense at all. This is consistent with a norm of equity, as it makes participation a revenue-neutral activity for all, no matter how much they earn. Nevertheless, it is very likely to reinforce unfair distribution of research risks and benefits between different social strata by prompting researchers to make savings by drawing research participants from “cheaper” populations, especially from the unemployed and the low-paid (Dickert and Grady 1999; Resnik 2015).

Thirdly, justice supports remuneration for participants’ contributions to the development of science and society, provided it is fair, i.e., equitable, adequate, and non-exploitative. Remuneration is equitable when it does not violate the norm “equal pay for equal work” (Dickert and Grady 1999; Resnik 2015; Gelinas et al. 2018, 2020; Persad et al. 2019). Equal time, efforts, burdens and risks associated with participation deserve equal remuneration (measured in market value, rather than in nominal value in case of multi-site studies conducted in different settings). However, when different groups of participants in a study are expected to make different contributions, equitable remuneration should reflect differences in their input (Persad et al. 2019). This is consistent with equal respect and concern for each and every individual and it prevents discrimination.

Remuneration is adequate when it is proportionate in value to the value of participants’ contribution to the study. The latter necessarily depends on various factors which determine how time- and effort-consuming, burdensome and risky give research project is. For example, when a study involves 2 visits, each lasting 60-min and requires filling a questionnaire and giving a blood sample for further analysis, and another project involves 4 visits of the same length and level of associated risks and burdens, participants of the latter study—ceteris paribus—should be offered remuneration of double the remuneration offered to the participant of the first project.

All guidelines and regulatory documents referred to above follow this normative logic by recommending or permitting paying research participants an “appropriate” or “proportionate” or “just and fair” remuneration for the time spent and other inconveniences resulting from the study participation (Council of Europe 2005b, sec. 64; Health Research Authority 2014, par. 3.4; Council for International Organizations of Medical Sciences 2016, p. 53; Food and Drug Administration 2018; National Health and Medical Research Council 2019, par. 1.1). Only a few, however, mention the level of research risks among factors which should be taken into consideration when determining an adequate amount of payment. Moreover, those that do so provide conflicting instructions on this matter. For example, the CIOMS Guidelines expressis verbis state that “the level of compensation should not be related to the level of risk that participants agree to undertake” (2016, Commentary on Guideline 13). In contract, the Australian National Health and Medical Research Council’s guideline reads: “In cases where risk may be considered as a factor in determining payment, payment of participants based on the degree of risk associated with the research is not prohibited, so long as there is evidence that a participant’s ability to provide valid consent is not likely to be compromised” (2019, par. 1.3.). Also the UK Health Research Authority “sympathises with the view that not to allow payments on the basis of risk would be unduly paternalistic in the absence of evidence that the participants’ ability to provide valid consent would be compromised” (2014, par. 3.1.).

These regulatory variations mirror the lack of consensus regarding risk-based payments among research ethicists (Grady 2001; Menikoff 2001; Jones and Liddell 2009; Saunders 2009; Grimwade et al. 2020; Lynch and Largent 2020; Jamrozik and Selgelid 2020). Although a detailed analysis of this issue goes beyond the scope of this analysis, three following arguments provide strong reasons for arguing that adequate remuneration for research subjects should be proportionate also to the level of risk involved in participation: (i) an argument from consistency, which notices that risk-based remuneration is accepted in many non-research contexts, e.g., in high-risk professions (Menikoff 2001; Jones and Liddell 2009); (ii) an argument from the nature of human “guinea pigging” which claims that the assumption of risk is often an essential contribution of research participants (Menikoff 2001; Równińska 2018; Malmqvist 2019); (iii) an argument for public trust, which argues that paying proportionally to the incurred risk meets expectations regarding the fair treatment of a significant portion of prospective subjects as well as researchers, REC/IRB members, and other members of the research community (Czarny et al. 2010; Ripley et al. 2010; Largent et al. 2012; Grimwade et al. 2020), thereby enhances public trust in research at large.

Finally, remuneration is non-exploitative when it is not lower than a socially accepted payment which is (or would be) offered for a similarly time- and effort-consuming, burdensome and risky activity, outside research context in the same setting (Gelinas et al. 2018; Largent and Lynch 2017b). Although space does not permit discussing in
details how a proper amount of such defined payment should be calculated, a proposal widely advocated in the literature should be mentioned here, namely a minimum hourly wage benchmark. Numerous authors and guidelines suggest that the amount should be based on the minimum hourly wage in the region or country as a point of reference (Council for International Organizations of Medical Sciences 2016, Commentary to Guideline 15, 53; National Health and Medical Research Council 2019, Appendix 1, 7) with augmentations for particularly burdensome procedures (Ackerman 1989; Dickert and Grady 1999; Grady 2005; Gelinas et al. 2020), risks involved (Menikoff 2001), and even other additional benefits (cf. Anderson and Weijer 2002). This proposal has three advantages. It is relatively easy to implement as it provides a clear method of setting the baseline amount of reimbursement for research participants. It allows for keeping payments sensitive to specific features of the project and subjects’ contributions by accepting relevant payment augmentations. And—since it sets the reference value of an non-exploitive remuneration relatively low—“more researchers could afford to pay a fair wage and fewer would be inconvenienced by a prohibition on unfairly low wages” (Phillips 2011a, p. 219). However, it also faces challenges. Offers of payment calculated in such a way would be attractive for prospective subjects with low SES, who have no reasonable alternatives to get engaged in better paid activities. But these offers would likely have no impact on recruitment of individuals who are better-off, i.e., who have better paid jobs or capacity to make more money outside research context, thus potentially biasing the subjects’ recruitment. In contrast, the average-wage benchmark for non-exploitive remuneration, mentioned by some authors (Phillips 2011a) would seem fairer as it would make payment offers reasonably attractive to both individuals of SES and to those better-off, thus promoting a fair distribution of research risks and benefits between different social strata.

Fourthly, the principle of justice does not prohibit remunerations going beyond what constitutes equitable, adequate, and minimally non-exploitive payment—i.e., remunerations driven by a market-driven forces. Although such payments are conceptualized as “price” for desired services, in order to avoid exploitation, they should not be lower than remuneration viewed as “reward”. They may, however, be higher than “rewards” and they may vary within the same study on the basis of salient characteristics of particular groups of participants (i.e., their age, sex, race, ethnicity, rare clinical status etc.). Offering disproportionally high and/or differential remuneration does not violate requirements of justice insofar as the payment is designed to help the study to meet its social and scientific goals by enhancing recruitment and retention of the necessary category of subjects, and it does not reproduce or reinforce wider social inequities and injustices, e.g., racial biases or class differences (Persad et al. 2019).

**Individual beneficence and payment**

The principle of individual beneficence is rarely invoked in the discussion on ethical payment practices. Most probably, it is because research is about advancing the interests of science and society, not the interests of individual participants (Emanuel et al. 2000; Miller and Brody 2003). While there are studies which have a potential of direct therapeutic benefits for the participants, it is neither the goal of research practice, nor a requirement that research should be beneficial for participants. Moreover, all canonical guidelines on human research ethics exclude non-direct and non-medical benefits to research subjects, such as payment, from the risk–benefit analysis, thereby forbidding IRBs/RECs to take payment into account as a benefit to counterbalance research risks (Emanuel et al. 2010; *contra* Wertheimer 2013).

Nevertheless, despite the above normative premises, in fact, individuals treat money as a benefit when considering an offer to participate, and deciding on entering the study (Abadie 2010; Czarny et al., 2010; Stunkel and Grady 2011; Grady et al. 2017; Fisher et al. 2018; Manton et al. 2019). For an individual prospective participant, the payment is a part of an equation for the overall attractiveness of a research project. Recompenses, when full and adequate, make the research participation cost-free for subjects, and—as shown above—they can remove at least some entrance barriers for patients to potentially clinically beneficial studies. Remunerations—whether calculated as a reward or as a price—may constitute a gain for subjects, thereby making participation in research overall beneficial from their personal perspective.

Thus, the principle of individual beneficence provides additional support for offering money to research subjects. However, the same principle justifies the claim that payment for participation should not be overly attractive. There is some evidence that too attractive remuneration may increase the risk of jeopardizing subjects’ health by encouraging them to conceal or misrepresent information important for their safety in order to ensure recruitment or continued participation in paid research (Bentley and Thacker 2004; Devine et al. 2013, 2015; Lee et al. 2018; Lynch et al. 2019). Such a deceptive behavior by participants may take various forms (e.g., nondisclosure of concurrent enrollment in other studies, concealment of tobacco use, alcohol consumption, or illicit substance abuse, concealment of pre-existing medical conditions, falsification of current health status, over-reporting of a study protocol adherence, etc.), and it may result in severe adverse events or even subjects’ death (Lee et al. 2018).
The ethical anatomy of payment for research participants

Table 1 The ethical anatomy of payment for research participants

| Principle of social beneficence | Subject to the reservations listed below, payment for research participants is prima facie obligatory as it promotes the realization of a common good (socially valuable knowledge). | Recompense for reasonable expenses | Recompense for loss of reasonably expected profits (e.g., wages) | Remuneration conceptualized as a reward for contribution (time, affords, burdens, risks) | Remuneration conceptualized as a price driven by supply-demand market forces |
|--------------------------------|-------------------------------------------------------------------------------------------------------------------|---------------------------------|---------------------------------------------------------------|----------------------------------------------------------------------------------|----------------------------------------------------------------------------------|
|                                | Subject to the reservations listed below, payment for research participants is prima facie obligatory as it promotes the realization of a common good (socially valuable knowledge). |       | Recompense for reasonable expenses | Recompense for loss of reasonably expected profits (e.g., wages) | Remuneration conceptualized as a reward for contribution (time, affords, burdens, risks) | Remuneration conceptualized as a price driven by supply-demand market forces |
| Principle of respect for autonomy | Subject to the reservations listed below, ethical payment promotes participants’ autonomy. | Enables realization of an individual’s free will to participate in research by removing economic entry barriers. | Enables realization of an individual’s free will to participate in research by removing economic entry barriers. | Enables realization of an individual’s free will to participate in research by removing economic entry barriers. | Enables realization of an individual’s free will to participate in research by removing economic entry barriers. |
| Principle of individual beneficence | Subject to the reservations listed below, ethical payment makes participation en bloc more beneficial to participants. | Reduces risk of participation having a negative impact on the subject’s economic position (welfare) by making participation a cost-free activity. | Reduces risk of participation having a negative impact on the subject’s economic position (welfare) by making participation a cost-free activity. | Reduces risk of participation having a negative impact on the subject’s economic position (welfare) by making participation a cost-free activity. | Reduces risk of participation having a negative impact on the subject’s economic position (welfare) by making participation a cost-free activity. |
| Principle of justice | Subject to the exclusion and reservations listed below, ethical payment promotes fair distribution of research risks and benefits. | Reduces inequality of opportunities to for individuals with low SES by removing economic entry barriers. | Reduces inequality of opportunities to for individuals with low SES by removing economic entry barriers. | Reduces inequality of opportunities to for individuals with low SES by removing economic entry barriers. | Reduces inequality of opportunities to for individuals with low SES by removing economic entry barriers. |

Conclusions

The paper argues that the ethical anatomy of paying research participants is built upon four basic principles of research ethics (and bioethics in general). The ethical “spine” of the practice is the principle of social beneficence, which requires the maximization of the common good—in the case of research practice—socially valuable scientific knowledge. This principle grounds a general prima facie moral obligation of offering payment to research subjects. The remaining ethical principles constitute a “skeleton” of morally sound payment practices by providing additional moral reasons for offering or not offering certain types of payments to research participants. As discussed above and presented in Table 1, the principles argue for offering research participants:

- recompense for reasonable expenses, but not for lost wages (or loss of other reasonably expected profits);
- remuneration conceptualized as a reward for their valuable contribution, provided (i) the remuneration meets standards of equality, adequacy and non-exploitation, and (ii) it is not overly attractive, i.e., it does not constitute undue inducement for participation or retention, and does not encourage deceptive behaviors;
- remuneration conceptualized as a market-driven price, provided (i) the remuneration is necessary and designed to help the study achieve its social and scientific goals, (ii) it does not reinforce wider social injustices and inequalities; (iii) it meets the requirement of non-exploitation; (iv) it is not overly attractive (as defined above).

Obviously, a proper application of this ethical “skeleton” into research practice requires investigators and RECs/IRBs to take into account “the nature of the study, the nature of participants contributions and vulnerabilities, institutional and organizational guidelines, and local and cultural norms” (Grady 2005, p. 1686). Moreover, to make the proposed scheme fully helpful in determining whether any particular offer of payment is not overly attractive and whether it meets standards of not equality, adequacy and non-exploitation, further detailed analyses of these standards are needed. The proposed scheme should also be tested against and enriched by further empirical studies about payment, especially about how money impacts subjects’ decision-making processes and behaviors.

Funding This analysis was supported by a grant of the National Science Centre, Poland, No. 2015/17/B/HS1/02390.
Data availability Not applicable.

Code availability Not applicable.

Declarations

Conflict of interest I have no conflict of interests or competing interests to disclose.

Open Access This article is licensed under a Creative Commons Attribution 4.0 International License, which permits use, sharing, adaptation, distribution and reproduction in any medium or format, as long as you give appropriate credit to the original author(s) and the source, provide a link to the Creative Commons licence, and indicate if changes were made. The images or other third party material in this article are included in the article’s Creative Commons licence, unless indicated otherwise in a credit line to the material. If material is not included in the article’s Creative Commons licence and your intended use is not permitted by statutory regulation or exceeds the permitted use, you will need to obtain permission directly from the copyright holder. To view a copy of this licence, visit http://creativecommons.org/licenses/by/4.0/. 

References

Abadie, Roberto. 2010. The professional guinea pig: Big pharma and the risky world of human subjects. Durham: Duke University Press.

Abadie, Roberto. 2015. The “mild-torture economy”: exploring the world of professional research subjects and its ethical implications. Physis: Revista De Saúde Coletiva 25: 709–728.

Ackerman, Terrence F. 1989. An ethical framework for the practice of paying research subjects. IRB: Ethics & Human Research 11 (4): 1–4.

Almeida, Luis, Benedita Azevedo, Teresa Nunes, Manuel Vaz-da-Silva, and Patricio Soares-da-Silva. 2007. Why healthy subjects volunteer for phase I studies and how they perceive their participation? European Journal of Clinical Pharmacology 63 (11): 1085–1094.

Altman, Douglas G. 1980. Statistics and ethics in medical research: III How large a sample? British Medical Journal 281 (6251): 1336–1338.

American Cancer Society Cancer Action Network. 2018. Barriers to patient enrollment in therapeutic clinical trials for cancer: a landscape report. https://www.acscan.org/policy-resources/clinical-trial-barriers. Accessed 28 August 2021.

Anderson, James A., and Charles Weijer. 2002. The research subject as wage earner. Theoretical Medicine and Bioethics 23 (4): 359–376.

Beauchamp, Tom L., and James F. Childress. 2001. The principles of biomedical ethics, 5th ed. Oxford: Oxford University Press.

Belfrage, Sara. 2016. Exploitative, irresistible, and coercive offers: Why research participants should be paid well or not at all. Journal of Global Ethics 12 (1): 69–86.

Bell, Jennifer A.H., and Lynda G. Balneaves. 2015. Cancer patient decision making related to clinical trial participation: An integrative review with implications for patients’ relational autonomy. Supportive Care in Cancer 23 (4): 1169–1196.

Bentley, John P., and Paul G. Thacker. 2004. The influence of risk and monetary payment on the research participation decision making process. Journal of Medical Ethics 30 (3): 293–298.

Bierer, Barbara E., Sarah A. White, Luke Gelinas, and David H. Strauss. 2021. Fair payment and just benefits to enhance diversity in clinical research. Journal of Clinical and Translational Science 5 (1): e159. https://doi.org/10.1017/cts.2021.816.

Caldwell, Patriona H., Sana Hamilton, Alvin Tan, and Jonathan C. Craig. 2010. Strategies for increasing recruitment to randomised controlled trials: Systematic review. PLoS Medicine 7 (11): e1000368.

Campbell, Marion K., Claire Snowdon, David Francis, Diana R. Elbourne, Alison M. McDonald, Rosemary C. Knight, Vikki Entwistle, Jo Garcia, Ian Roberts, and STEPS Group. 2007. Recruitment to randomised trials: Strategies for trial enrolment and participation study. The STEPS study. Health Technology Assessment. https://www.journalslibrary.nihr.ac.uk/hta/hta11480/#/full-report. Accessed 28 Aug 2021.

Carlisle, Benjamin, Jonathan Kimmelman, Tim Ramsay, and Nathalie MacKinnon. 2015. Unsuccessful trial accrual and human subject protections: An empirical analysis of recently closed trials. Clinical Trials 12 (1): 77–83.

Center for Information and Study on Clinical Research Participation. 2019. Charts and statistics. https://www.ciscrp.org/education-center/charts-statistics. Accessed 28 Aug 2021.

Chams, Tod. 2001. Participation as commodity, participation as gift. American Journal of Bioethics 1 (2): 48–48.

Chino, Fumiko, and S Yousuf Zafar. 2019. Financial toxicity and equitable access to clinical trials. American Society of Clinical Oncology Educational Book 39: 11–18.

Cooper, Melinda, and Catherine Waldby. 2014. Clinical labor: tissue donors and research subjects in the global bioeconomy. Durham: Duke University Press.

Council for International Organizations of Medical Sciences. 2016. International ethical guidelines for health-related research involving humans. Geneva: Council for International Organizations of Medical Sciences. https://cioms.ch/wp-content/uploads/2017/01/WEB-CIOMS-EthicalGuidelines.pdf. Accessed 28 August 2021.

Council of Europe. 2005a. Additional Protocol to Convention on human rights and biomedicine concerning biomedical research. CETS 195. Strasbourg: Council of Europe. https://rm.coe.int/168008371a. Accessed 28 Aug 2021.

Council of Europe. 2005b. Explanatory report to Additional Protocol to Convention on human rights and biomedicine concerning biomedical research. Strasbourg: Council of Europe. https://rm.coe.int/16800d3810. Accessed 28 Aug 2021.

Cryder, Cynthia E., Alex J. London, Kevin G. Volpp, and George Loewenstein. 2010. Informative inducement: Study payment as a signal of risk. Social Science & Medicine 70 (3): 455–464.

Czarny, Matthew J., Nancy E. Kass, Charles W. Flexner, Kathryn A. Car- son, Rachel K. Myers, and Ephraim J. Fuchs. 2010. Payment to healthy volunteers in clinical research: The research subject’s perspective. Clinical Pharmacology & Therapeutics 87 (3): 286–293.

Department of Health and Human Services. 2018. U.S. Code of Federal Regulations, Title 45: Public Welfare, Part 46: Protection of human subjects. https://www.hhs.gov/ohrp/regulations-and-policies/regulations/45-cfr-46/revised-common-rule-regulatory-text/index.html. Accessed 28 Aug 2021.

Devine, Eric G., Megan E. Waters, Megan Putnam, Caitlin Surprise, Katie O’Malley, Courtney Richambault, Rachel L. Fishmana, Clifford M. Knappa, Elissa H. Pattersona, Ofra Sarid-Segala, Chris Streetera, Laurie Colanaria, and Domenic A. Ciraulo. 2013. Concealment and fabrication by experienced research subjects. Clinical Trials 10 (6): 935–948.

Devine, Eric G., Clifford M. Knapp, Ofra Sarid-Segala, Sean M. O’Keefe, Cale Wardell, Morgan Baskett, Ashley Pecccia, Katie Ferrell, and Domenic A. Ciraulo. 2015. Payment expectations for research participation among subjects who tell the truth, subjects who conceal information, and subjects who fabricate information. Contemporary Clinical Trials 41: 55–61.
Dickert, Neal W. 2009. Enrollment of economically disadvantaged participants in clinical research. *AMA Journal of Ethics* 11 (1): 54–60.

Dickert, Neal W. 2013. Concealment and fabrication: The hidden price of payment for research participation? *Clinical Trials* 10 (6): 840–841.

Dickert, Neal, Ezekiel Emanuel, and Christine Grady. 2002. Paying research subjects: An analysis of current policies. *Annals of Internal Medicine* 136 (5): 368–373.

Dickert, Neal, and Christine Grady. 1999. What’s the price of a research subject? Approaches to payment for research participation. *The New England Journal of Medicine* 341: 198–203.

Dickert, Neal, and Christine Grady. 2008. Incentives for participants. In *The Oxford textbook on the ethics of clinical research*, ed. Ezekiel J. Emanuel, Robert A. Crouch, Christine C. Grady, Reidar K. Lie, Franklin G. Miller, and David D. Wendler, 386–396. New York: Oxford University Press.

Dresser, Rebecca. 2013. Subversive subjects: Rule-breaking and deception in clinical trials. *The Journal of Law, Medicine & Ethics* 41 (4): 829–840.

Duncan, Peter. 2010. Health, health care and the problem of intrinsic value. *Journal of Evaluation in Clinical Practice* 16 (2): 318–322.

Dunn, Laura B., and Nora E. Gordon. 2005. Improving informed consent and enhancing recruitment for research by understanding economic behavior. *JAMA* 293 (5): 609–612.

Elliott, Carl. 2008. Guinea-pigging. *The New Yorker*, January 7, pp. 36–41.

Elliott, Carl, and Roberto Abadie. 2008. Exploiting a research under-class in phase 1 clinical trials. *The New England Journal of Medicine* 358 (22): 2316–2317.

Emanuel, Ezekiel J. 2004. Ending concerns about undue inducement. *Journal of Law, Medicine & Ethics* 32 (1): 100–105.

Emanuel, Ezekiel J. 2005. Undue inducement: Nonsense on stilts? *The American Journal of Bioethics* 5 (5): 9–13.

Emanuel, Ezekiel J., David Wendler, and Christine Grady. 2000. What makes clinical research ethical? *JAMA* 283 (20): 2701–2711.

European Union. 2014. Regulations (EU) No 536/2014 of the European Parliament and of the Council of 16 April 2014 on clinical trials on medicinal products for human use, and repealing Directive 2001/20/EC. https://ec.europa.eu/health/sites/default/files/files/eudralex/vol-1/reg_2014_536/reg_2014_536_en.pdf. Accessed 28 Aug 2021.

Faden, Ruth, and Tom Beauchamp. 1986. *A history and theory of informed consent*. New York: Oxford University Press.

Fisher, Jill A. 2019. Research payment and its social justice concerns. *The American Journal of Bioethics* 19 (9): 35–36.

Fisher, Jill A., Lisa McManus, Megan M. Wood, Marci D. Cottingham, Julianne M. Kalbaugh, Torin Monahan, and Rebecca L. Walker. 2018. Healthy volunteers’ perceptions of the benefits of their participation in phase I clinical trials. *Journal of Empirical Research on Human Research Ethics* 13 (5): 494–510.

Fisher, Jill A., Torin Monahan, and Rebecca L. Walker. 2019. Picking and choosing among phase I trials. *Journal of Bioethical Inquiry* 16 (4): 535–549.

Food and Drug Administration. 2018. Payment and reimbursement to research subjects. Guidance for institutional review boards and clinical investigators. U.S. Food and Drug Administration: Office of the Commissioner, Office of Clinical Policy and Programs, Office of Clinical Policy, Office of Good Clinical Practice. https://www.fda.gov/regulatory-information/search-fda-guidance-documents/payment-and-reimbursement-research-subjects. Accessed 28 Aug 2021.

Friedman, Daniela B., Caroline Foster, Caroline D. Bergeron, Andrea Tanner, and Sei-Hill. Kim. 2015. A qualitative study of recruitment barriers, motivators, and community-based strategies for increasing clinical trials participation among rural and urban populations. *American Journal of Health Promotion* 29 (5): 332–338.

Fry, Craig L., Alison Ritter, Simon Baldwin, Kathryn J. Bowen, T. Paul Gardiner, Rebecca Jenkinson Holt, and Jennifer Johnston. 2005. Paying research participants: A study of current practices in Australia. *Journal of Medical Ethics* 31 (9): 542–547.

Gelinas, Luke, Emily A. Largent, I. Glenn Cohen, Susan Kornetsky, Barbara E. Bierer, and Holly Fernandez-Lynch. 2018. A framework for ethical payment to research participants. *The New England Journal of Medicine* 378 (8): 766–771.

Gelinas, Luke, Sarah A. White, and Barbara E. Bierer. 2020. Economic vulnerability and payment for research participation. *Clinical Trials* 17 (3): 264–272.

Geller, Stacie E., Abby Koch, Beth Pellettieri, and Molly Carnes. 2011. Inclusion, analysis, and reporting of sex and race/ethnicity in clinical trials: Have we made progress? *Journal of Women’s Health* 20 (3): 315–320.

Glannon, William. 2006. Phase I oncology trials: Why the therapeutic misconception will not go away. *Journal of Medical Ethics* 32: 252–255.

Grady, Christine. 2001. Money for research participation: Does it jeopardize informed consent? *The American Journal of Bioethics* 1 (2): 40–44.

Grady, Christine. 2005. Payment of clinical research subjects. *The Journal of Clinical Investigation* 115 (7): 1681–1687.

Grady, Christine, Gabriella Bedarid, Ninet Sinaii, Mark A. Gregorio, and Ezekiel J. Emanuel. 2017. Motivations, enrollment decisions, and socio-demographic characteristics of healthy volunteers in phase 1 research. *Clinical Trials* 14 (5): 526–536.

Grady, Christine, Neal Dickert, Tom Jawetz, Gary Gensler, and Ezekiel J. Emanuel. 2005. An analysis of US practices of paying research participants. *Contemporary Clinical Trials* 26 (3): 365–375.

Grant, Ruth W., and Jeremy Sugarman. 2004. Ethics in human subjects research: Do incentives matter? *The Journal of Medicine and Philosophy* 29 (6): 717–738.

Grimwade, Olivia, Julan Savulescu, Alberto Giubilini, Justin Oakley, Joshua Osowicki, Andrew J. Pollard, and Anne-Marie Nussberger. 2020. Payment in challenge studies: Ethics, attitudes and a new payment for risk model. *Journal of Medical Ethics* 46 (12): 815–826.

Gul, Raisa B., and Parveen A. Ali. 2010. Clinical trials: The challenge of recruitment and retention of participants. *Journal of Clinical Nursing* 19 (1–2): 227–233.

Halpern, Scott D., Jason H.T. Karlwaisch, and Jesse A. Berlin. 2002. The continuing unethical conduct of underpowered clinical trials. *JAMA* 288 (3): 358–362.

Hamel, Lauren M., Louis A. Penner, Terrance L. Albrecht, Elisabeth Heath, Klement C. Gwede, and Susan Eggle. 2016. Barriers to clinical trial enrollment in racial and ethnic minority patients with cancer. *Cancer Control* 23 (4): 327–337.

Health Research Authority. 2014. Ethics guidance: payments and incentives in research UK NHS Health Research Authority. National Research and Ethics Advisors’ Panel. https://s3.eu-west-2.amazonaws.com/www.hra.nhs.uk/media/documents/hra-guidance-documents/payment-and-reimbursement-research-subjects. Accessed 28 Aug 2021.

Heller, Caren, Joyce E. Balls-Berry, Jill D. Nery, Patricia J. Erwin, Dawn Littleton, Mimi Kim, and Winston P. Kuo. 2014. Strategies addressing barriers to clinical trial enrollment of underrepresented populations: A systematic review. *Contemporary Clinical Trials* 39 (2): 169–182.

International Conference on Harmonisation of Technical Requirements for Registration of Pharmaceuticals for Human Use. 2016. ICH Harmonised Guideline Integrated Addendum To ICH E6(R1): Guideline For Good Clinical Practice E6(R2). https://database.
Lemmens, Trudo, and Carl Elliott. 1999. Guinea pigs on the pay roll: The ethics of paying research subjects. *Accountability in Research* 7 (1): 3–20.

Lemmens, Trudo, and Carl Elliott. 2001. Justice for the professional guinea pig. *American Journal of Bioethics* 1 (2): 51–53.

London, Alex J. 2003. Threats to the common good: Biochemical weapons and human subjects research. *Hastings Center Report* 33 (5): 17–25.

London, Alex J. 2006. Reasonable risks in clinical research: A critique and a proposal for the Integrative Approach. *Statistics in Medicine* 25 (17): 2869–2885.

London, Alex J., Jonathan Kimmelman, and Marina E. Emborg. 2010. Beyond access v. protection in trials of innovative therapies. *Science* 328: 829–830.

Lynch, Holly F. 2014. Human research subjects as human research workers. *Yale Journal of Health Policy, Law, and Ethics* 14: 122–193.

Lynch, Holly F., Thomas C. Darton, Jae Levy, Frank McCormick, Ubaka Ogbugo, Puth O. Payne, Alvin E. Roth, Akiiah Jefferson Shah, Thomas Smiley, and Emily A. Largent. 2021. Promoting ethical payment in human infection challenge studies. *The American Journal of Bioethics* 21 (3): 11–31.

Lynch, Holly F., Steven Joffe, Harsha Thirumurthy, Dawei Xie, and Emily A. Largent. 2019. Association between financial incentives and participant deception about study eligibility. *JAMA Network Open* 2 (1): e187355. https://doi.org/10.1001/jamanetworkopen.2018.7355.

Lynch, Holly F., and Emily A. Largent. 2020. Compensating for research risk: Permissible but not obligatory. *Journal of Medical Ethics* 46 (12): 827–828.

MacKay, Douglas, and Rebecca L. Walker. 2021. Paying for fairness? Incentives and fair subject selection. *The American Journal of Bioethics* 21 (3): 35–37.

Macklin, Ruth. 1981. ‘Due’ and ‘undue’ inducements: on paying money to research subjects. *IRB: Ethics & Human Research* 3 (5): 1–6.

Macklin, Ruth. 1989. The paradoxical case of payment as benefit to research subjects. *IRB: Ethics & Human Research* 11 (6): 1–3.

Malmqvist, Erik. 2019. “Paid to endure”: Paid research participation, passivity, and the goods of work. *The American Journal of Bioethics* 19 (9): 11–20.

Manton, Kerry J., Cassandra S. Gould, Katherine M. White, Paul M. Griffin, and Suzanne L. Elliott. 2019. Qualitative study investigating the underlying motivations of healthy participants in phase I clinical trials. *British Medical Journal Open* 9: e024224. https://doi.org/10.1136/bmjopen-2018-024224.

McCann, Sharon K., Marion K. Campbell, and Vikki A. Entwistle. 2010. Reasons for participating in randomised controlled trials: Conditional altruism and considerations for self. *Trials* 11 (1): 1–10.

McManus, Lisa, and Jill A. Fisher. 2018. To report or not to report: Exploring healthy volunteers’ rationales for disclosing adverse events in Phase I drug trials. *AJOB Empirical Bioethics* 9 (2): 82–90.

McNeill, Paul. 1997. Paying people to participate in research: Why not? *Bioethics* 11 (5): 390–396.

Menikoff, Jerry. 2001. Just compensation: Paying research subjects relative to the risks they bear. *The American Journal of Bioethics* 1 (2): 56–58.

Miller, Franklin G., and Howard Brody. 2003. A critique of clinical equipoise: Therapeutic misconception in the ethics of clinical trials. *The Hastings Center Report* 33 (3): 19–28.

Millum, Joseph, and Michael Garnett. 2019. How payment for participation can be coercive. *The American Journal of Bioethics* 19 (9): 21–31.

Moriarty, Jeffrey. 2020. What’s in a wage? A new approach to the justification of pay. *Business Ethics Quarterly* 30 (1): 119–137.

National Commission for the Protection of Human Subjects of Biomedical and Behavioral Research. 1979. *The Belmont report: Ethical principles and guidelines for the protection of human subjects of research*. U.S. Department of Health and Human Services. https://www.hhs.gov/ohrp/regulations-and-policy/belmont-report/read-the-belmont-report/index.html. Accessed 28 Aug 2021.

National Health and Medical Research Council. 2019. *Payment of participation in research: information for researchers*, *HRECs and other ethics review bodies*. Australian Research Council and Universities Australia. Commonwealth of Australia, Canberra. https://www.nhmrc.gov.au/about-us/publications/payment-participants-research-information-researchers-hrecs-and-other-ethics-review-bodies#block-views-block-file-attachments-content-block-1. Accessed 28 Aug 2021.

Ndebele, Paul, Joseph Mufutso-Bengo, and Takaﬁra Mduluza. 2008. Compensating clinical trial participants from limited resource settings in internationally sponsored clinical trials: A proposal. *Malawi Medical Journal* 20 (2): 42–45.
The ethical anatomy of payment for research participants

Nipp, Ryan D., Kesselyong, and Electra D. Paskett. 2019a. Overcoming barriers to clinical trial enrollment. American Society of Clinical Oncology Educational Book 39: 105–114.

Nipp, Ryan D., Hang Lee, Emily Gorton, Morgan Lichtenstein, Salome Kuchukhidze, Elyse Park, Bruce A. Chabner, and Beverly Moy. 2019b. Addressing the financial burden of cancer clinical trial participation: Longitudinal effects of an equity intervention. The Oncologist 24 (8): 1048–1055.

Nuffield Council on Bioethics. 2011. Human bodies: donation for medicine and research. Nuffield Council on Bioethics. https://www.nuffieldbioethics.org/publications/human-bodies-donation-for-medicine-and-research. Accessed 28 Aug 2021.

Nussbaum, Martha C. 2015. Creating capabilities: The human development approach. Cambridge: Harvard University Press.

Parkinson, Beth, Rachel Meercock, Matt Sutton, Eleonora Fichera, Nicola Mills, Gillian W. Shorter, S. Treweek, Nicola L. Harman, Rebecca P. Brown, Katie Giles, and Peter Bower. 2019. Designing and using incentives to support recruitment and retention in clinical trials: A scoping review and a checklist for design. Trials 20 (1): 1–14.

Pasqualetti, Giuseppe, Giovanni Gori, Corrado Blandizzi, and Mario Del Taccia. 2010. Healthy volunteers and early phases of clinical experimentation. European Journal of Clinical Pharmacology 66 (7): 647–653.

Persad, Govind, Holly Fernandez Lynch, and Emily Largent. 2019. Differential payment to research participants in the same study: An ethical analysis. Journal of Medical Ethics 45 (5): 318–322.

Phillips, Trisha. 2011a. Exploitation in payments to research subjects. Bioethics 25 (4): 209–219.

Phillips, Trisha. 2011b. A living wage for research subjects. The Journal of Law, Medicine & Ethics 39 (2): 243–253.

Probstfield, Jeffrey L., and Robert L. Frye. 2011. Strategies for recruitment and retention of participants in clinical trials. JAMA 306 (6): 1798–1799.

Radder, Hans. 2017. Which scientific knowledge is a common good? Social Epistemology 31 (5): 431–450.

Rawls, John. 1971. A theory of justice. Cambridge: Harvard University Press.

Resnik, David B. 2011. Scientific research and the public trust. Science and Engineering Ethics 17 (3): 399–409.

Resnik, David B. 2015. Bioethical issues in providing financial incentives to research participants. Medicolegal and Bioethics 5: 35–41.

Resnik, David B. 2019. Are payments to human research subjects ethical or suspect? Journal of Clinical Research Best Practices 15 (6): 2374.

Rhodes, Rosamond. 2010. Rethinking research ethics. The American Journal of Bioethics 10 (19): 10–36.

Ripley, Elizabeth, Francis Macrina, Monika Markowitz, and Chris Gennings. 2010. Why do we pay? A national survey of investigators and IRB chairpersons. Journal of Empirical Research onInvestigators and IRB chairpersons. Journal of Empirical Research on Human Research Ethics 3 (5): 43–56.

Roche, Eric, Romain King, Helen M. Mohan, Blanaid Gavin, and Fiona McNicholas. 2013. Payment of research participants: Current practice and policies of Irish research ethics committees. Journal of Medical Ethics 39 (9): 591–593.

Różyńska, Joanna. 2015. On the alleged right to participate in high-risk research. Bioethics 29 (7): 451–461.

Różyńska, Joanna. 2018. What makes clinical labour different? The case of human guinea pigging. Journal of Medical Ethics 44 (9): 638–642.

Różyńska, Joanna. 2021. Research participants should be rewarded rather than “compensated for time and burdens.” The American Journal of Bioethics 21 (3): 53–55.

Salman, Rustam A. S. Elaine Beller, Jonathan Kagan, Elina Hemminki, Robert S. Phillips, Julian Savulescu, Malcolm Macleod, Janet Wisley, and Iain Chalmers. 2014. Increasing value and reducing waste in biomedical research regulation and management. The Lancet 383 (9912): 176–185.

Saunders, John. 2009. Should healthy volunteers in clinical trials be paid according to risk? BMJ. https://doi.org/10.1136/bmj.b4145.

Schaefer, G. Owen., Ezekiel J. Emanuel, and Alan Wertheimer. 2009. The obligation to participate in biomedical research. JAMA 302 (1): 67–72.

Shamoo, Adi E., and David B. Resnik. 2006. Strategies to minimize risks and exploitation in phase one trials on healthy subjects. The American Journal of Bioethics 6 (3): W1–W13.

Stones, Martyn, and John McMillan. 2010. Payment for participation in research: a pursuit for the poor? Journal of Medical Ethics 36 (1): 34–36.

Stunkel, Leanne, and Christine Grady. 2011. More than the money: A review of the literature examining healthy volunteer motivations. Contemporary Clinical Trials 32 (3): 342–352.

Sully Ben, G., Steven A. Julious, and Jon A. Nicholl. 2013. A reinvestigation of recruitment to randomised, controlled, multicenter trials: A review of trials funded by two UK funding agencies. Trials 14: 166.

Tishler, Carl L., and Suzanne Bartholomae. 2002. The recruitment of normal healthy volunteers: A review of the literature on the use of financial incentives. The Journal of Clinical Pharmacology 42 (4): 365–375.

Treweek, Shaun, Pauline Lockhart, Marie Pitkethly, Jonathan A. Cook, Monica Kjeldstrøm, Matir Jihotansen, Taina K. Taskila, Frank M. Sullivan, Sue Wilson, Catherine Jackson, Ritu Jones, and Elizabeth D. Mitchell. 2013. Methods to improve recruitment to randomised controlled trials: Cochrane systematic review and metaanalysis. British Medical Journal Open: e002360. https://doi.org/10.1136/bmjopen-2012-002360.

Unger, Joseph M., Julie R. Gralow, Kathy S. Albain, Scott D. Ramsey, and Dawn L. Hershman. 2016. Patient income level and cancer clinical trial participation: A prospective survey study. JAMA Oncology 2 (1): 137–139.

Unger, Joseph M., Dawn L. Hershman, Kathy S. Albain, Carol M. Moinpour, Judith A. Petersen, Kenda Burg, and John J. Crowley. 2013. Patient income level and cancer clinical trial participation. Journal of Clinical Oncology 31 (5): 536–542.

Unger, Joseph M., Riha Vaidya, Dawn L. Hershman, Lori M. Minasian, and Mark E. Fleury. 2019. Systematic review and meta-analysis of the magnitude of structural, clinical, and physician and patient barriers to cancer clinical trial participation. JNCI: Journal of the National Cancer Institute 111 (3): 245–255.

Walker, Rebecca L., and Jill A. Fisher. 2019. “My body is one of the best commodities”: Exploring the ethics of commodification in phase 1 healthy volunteer clinical trials. Kennedy Institute of Ethics Journal 29 (4): 305–331.

Watson, Judith M., and David J. Torgerson. 2006. Increasing recruitment to randomised trials: A review of randomised controlled trials. BMC Medical Research Methodology 6 (1): 1–9.

Wertheimer, Alan. 2008. Exploitation in clinical research. In The Oxford textbook on the ethics of clinical research, ed. Ezekiel J. Emanuel, Robert A. Crouch, Christine C. Grady, Reidar K. Lie, Franklin G. Miller, and David D. Wendler, 201–210. New York: Oxford University Press.

Wertheimer, Alan. 2013. Is payment a benefit. Bioethics 27: 105–116.

Wertheimer, Alan, and Franklin G. Miller. 2008. Payment for research participation: A coercive offer? Journal of Medical Ethics 34 (5): 389–392.

Wilkinson, Martin, and Andrew Moore. 1997. Inducement in research. Bioethics 11 (5): 373–389.

Williams, Rebecca J., Tony Tse, Katelyn DiPiazza, and Deborah A. Zarins. 2015. Terminated trials in the ClinicalTrials.gov results database: Evaluation of availability of primary outcome data and reasons for termination. PLoS ONE 10 (5): e0127242.
Winter, Stuart S., Janet M. Page-Reeves, Kimberly A. Page, Emily Haozous, Angelica Solares, Carla N. Cordova, and Richard S. Larson. 2018. Inclusion of special populations in clinical research: Important considerations and guidelines. *Journal of Clinical and Translational Research* 4 (1): 56–69.

World Medical Association (WMA). 2013. Declaration of Helsinki. Ethical principles for medical research involving human subjects. Fortaleza, Brazil. https://www.wma.net/policies-post/wma-declaration-of-helsinki-ethical-principles-for-medical-research-involving-human-subjects/. Accessed 28 Aug 2021.

**Publisher's Note** Springer Nature remains neutral with regard to jurisdictional claims in published maps and institutional affiliations.