Perception of clinical research among patients and healthy volunteers of clinical trials

Felix Bergmann1,2 · Peter Matzneller1,3 · Maria Weber1 · Lusine Yeghiazaryan4 · Thorsten Fuereder5 · Thomas Weber6 · Markus Zeitlinger1

Received: 21 March 2022 / Accepted: 18 July 2022 / Published online: 27 July 2022 © The Author(s) 2022

Abstract
Purpose Clinical research relies on data from patients and volunteers, yet the target sample size is often not achieved. Here, we assessed the perception of clinical research among clinical trial participants to improve the recruitment process for future studies.

Methods We conducted a single-center descriptive and exploratory study of 300 current or former participants in various phase I–III clinical trials. Questionnaires were either distributed to current clinical trial participants or emailed to former subjects.

Results Subjects strongly agreed or agreed that contributing to improving medical care (> 81%), contributing to scientific research (> 79%), and trusting their treating physicians (> 77%) were motives for study participation. Among healthy volunteers, financial motives positively correlated with the number of clinical trials they had participated in (p < 0.05). Higher age positively correlated with expectation of best available treatment during study participation among patients (p < 0.05). Less than 8% of all subjects expressed “great concern” about the potential risks of sharing their personal information as part of the study. Subjects displayed “great trust” or “trust” in medical staff (86.6%) and in government research institutions (76.4%), and “very little trust” or “little trust” in pharmaceutical companies (35.4%) and health insurance companies (16.9%).

Conclusion Altruistic motives and trust in treating physicians were predominant motives for clinical trial participation. Older patients expected to receive the best available treatment during participation. Healthy volunteers who reported financial motives had participated in more clinical trials. Consistent with great trust in medical staff and government research institutions, little concern was expressed about the misuse of personal data during the trial.

Keywords Clinical research · Treating physicians · Volunteers

Introduction

Unbiased clinical safety and efficacy assessments of novel therapeutic approaches largely rely on randomized controlled trials (RCTs) [1–3]. However, the inability to recruit and retain sufficient numbers of participants remains a common problem in achieving conclusive results [4–8]. This may result in underpowered clinical trials, in which clinically relevant differences are reported as statistically insignificant [9]. Thus, several studies have investigated influential factors of trial recruitment and participation. Key influential factors include altruistic motives [10–14], such as contribution to

© Markus Zeitlinger
Markus.zeitlinger@meduniwien.ac.at
1 Department of Clinical Pharmacology, Medical University of Vienna, Währinger Gürtel 18-20, 1090 Vienna, Austria
2 Clinical Division of Plastic and Reconstructive Surgery, Department of Surgery, Medical University of Vienna, Währinger Gürtel 18-20, 1090 Vienna, Austria
3 Service of Rheumatology, Hospital of Merano (SABES-ASDAA), Via Rossini 5, 39012 Merano, Italy
4 Center for Medical Statistics, Informatics, and Intelligent Systems, Medical University of Vienna, Währinger Gürtel 18-20, 1090 Vienna, Austria
5 Division of Oncology, Department of Medicine I, Medical University of Vienna, Währinger Gürtel 18-20, 1090 Vienna, Austria
6 Division of Gastroenterology and Hepatology, Department of Medicine III, Medical University of Vienna, Währinger Gürtel 18-20, 1090 Vienna, Austria
scientific research, financial incentives [13, 15–18], and the doctor-patient relationship [11, 19, 20].

Another aspect closely related to clinical research is the handling of personal data collected during clinical trials. The European Medicines Agency (EMA) argues that access to adequately anonymized data collected from clinical trials may benefit the process of drug development [21]. Several studies highlight the potential benefits of data sharing, such as the acceleration of scientific discovery, improvement of cost-effectiveness, enabling comparisons between different populations, improving surveillance of drug safety and efficacy, increasing sample size, improvement of comparator-effectiveness analysis, and reduction of duplicated efforts [21–23]. Although hesitancy and concern have been expressed about widespread data sharing [24, 25], a previous study demonstrated that most clinical trial participants believe that the potential benefits outweigh the risks [14].

Furthermore, certain personality traits have been correlated with motives for clinical trial participation [26]. The Big Five personality traits (BIFI), also known as the OCEAN model, use survey data to describe aspects of personality [27]. The theory proposes the following five dimensions to describe the human psyche: Openness, Conscientiousness, Extraversion, Agreeableness, and Neuroticism [28]. However, when conducting clinical trials, time is often limited, and personality may not be of primary interest. In this case, the Ten-Item Personality Inventory (TIPI) offers an abbreviated (2-measure) alternative, when brevity is of importance [29, 30].

In light of these findings, the aim of this study was to further assess clinical trial participants’ motives for trial participation and their views on the risks of data sharing and to investigate personality as a predictor of clinical trial participation behavior.

Methods

Trial design

We conducted a single-center descriptive and exploratory study at the General Hospital, Vienna, Austria.

Participants

The patient collective of the study consisted of current or former participants of phase I–III clinical studies conducted at the Medical University of Vienna. All participants underwent an informed consent process and were enrolled between March 2019 and September 2020.

To generate a diverse sample of clinical trials, two identical adaptations of the questionnaire were created, one in physical print, the other as a digital survey. Printed surveys were distributed at 5 different departments to their clinical trial patients. Patients either completed the form immediately or returned it later by mail. The digital survey was distributed via email to patients or subjects who had either participated in a clinical trial in the past or were currently participating in one and had agreed to receive information about future trials.

Questionnaire

A 50-item questionnaire was designed to capture key parameters reflecting participants’ views on the recruitment process, motives for participation, trust in fields of scientific research, and their opinions on clinical trial data sharing with respect to their most recent trial participation. It also included questions regarding the participant’s demographics and assessed personality traits based on the TIPI. The survey was designed under consultation of the Statistics Department of the Medical University of Vienna and underwent further refinement after a pilot phase with 5 participants. The full text of the study questionnaire can be found in the Supplementary Appendix.

Ten-item personality inventory

The TIPI consists of 10 short personality questions [29]. It incorporates 2 questions to evaluate each trait of the OCEAN model. Answers were recorded on a Likert scale ranging from 1 to 5 (1 = Strongly disagree, 5 = Strongly agree). One of the corresponding questions in each case functions as a negative control and must be inverted (i.e., 1 is recoded to 5, 2 is recoded to 4). Finally, the mean of both answers correlating to their respective trait is assessed.

Statistical analysis

Statistical analysis was performed using version 4.0.3 of the commercially available computer program R. Bar charts and frequency tables were created for each categorical variable. Spearman correlation coefficients were calculated between the variables. Correlation tests were also performed to determine significance. The significance level \( \alpha \) was set at 0.05. All percentages respectively refer to the total number of responses obtained for each question.

Results

Study population

A total of 300 participants were included in the study (Fig. 1). Of the 3020 emails extracted from the database, 2430 emails were successfully sent (80.5%). Message delivery failures were received from the remaining 590 emails.
Two hundred sixty completed surveys (10.7%) were returned from the 2430 invited trial participants. The number of emails successfully sent, yet not opened (e.g., marked as spam or unread) could not be determined. A total of 40 of the 70 printed questionnaires (57.1%) were returned. Table 1 describes the characteristics of the sample.

**Process of informed consent**

Our results show that most participants recalled undergoing an elaborate process of informed consent with a typical duration of 5–15 min (65.6%). Goals, execution, and time expenditure of the study were most often recalled (> 90%), whereas information regarding personal rights, data protection, and anonymity were often reported as missing (recalled by 59.9%, 63.9%, and 68%, respectively) (Fig. S1). The majority of participants were either approached by the study doctor (39.8%) or their attending physician (19.7%) and reported easily comprehensible study goals (86.9% very good comprehensibility or good comprehensibility) (Figs. S2 and S3). Subjects were most often contacted in person (38.4%), via telephone (34%), or email (19.4%) (Fig. S4).

**Motives**

**Patients**

Predominant motives for trial participation among patients were trust in the attending physician, improving future medical care, and contribution to research (94.1%, 86.9%, and 81.9% strongly agreed or agreed, respectively) (Fig. 2). The majority of patients also strongly agreed or agreed that they expected best possible medical treatment (74.7%), closer medical supervision (60.3%), and better quality of care (51.3%). Patients did not express substantial concern that their illness would worsen, if they did not participate in the trial (58.3% strongly disagreed or disagreed). Increasing age positively correlated with expecting the best available treatment ($p < 0.05$) (Fig. S5).

**Healthy volunteers**

56.3% of healthy volunteers strongly agreed or agreed to participate based on financial incentives, outweighed by altruistic motives such as contribution to improvement of medical care and scientific research (81.6% and 79.7%, respectively)
and trust in their attending doctor (77.6%). However, only the indication of financial motives for participation positively correlated with the number of studies subjects had participated in ($p < 0.05$) (Fig. S6). Among healthy volunteers, female sex and increasing age positively correlated with the motives, “contribution to scientific research” ($p < 0.05$) and negatively correlated with financial motivation ($p < 0.05$). Increasing age also positively correlated with the motive “improvement of care for future patients” ($p < 0.05$). Higher income negatively correlated with financial incentives and the motive of trusting their attending doctor ($p < 0.05$).

**Expected beneficiaries of clinical trials**

More than 80% of all respondents expected companies that develop drugs and medicinal products (83.7%), researchers (82.1%), and patients (80.9%) to “greatly benefit” or “benefit” from clinical trials (Fig. 3). 63.1% of subjects had this expectation for doctors that treat patients.

**Data protection**

Less than 10% of all respondents said they were “very concerned” and less than one-third were “concerned” or “moderately concerned” about the potential consequences of sharing personal data (Fig. 4). Respondents were most concerned that their data could be used for marketing purposes instead of scientific purposes (20% “very concerned” or “concerned”) or that companies or individuals could make a great economic profit by using their data (17.2%). Participants were least concerned that they could experience personal disadvantages (8.8% “very concerned” or “concerned”) or that their data could be traceable with good computer knowledge (12.8%). Greater trust in scientific research correlated with lower concern regarding data sharing (Fig. S7). No disparity was observed when comparing healthy volunteers and patients, independent of gender and age ($p > 0.05$, data not shown).

**Trust in fields of scientific research**

Figure 5 documents participants’ levels of trust in fields of scientific research. Strong majorities of respondents expressed “great trust” or “trust” in medical staff (86.6%) and in government research institutions (76.4%); however, they showed “very little trust” or “little trust” in pharmaceutical companies (35.5%) and health insurance companies (16.9%). Trust in pharmaceutical companies, was higher among patients than healthy volunteers ($p < 0.05$) and increased with higher age among all individuals ($p < 0.05$).

**Ten-item personality index**

All pairs of questions that corresponded to a personality trait correlated indirectly ($p < 0.05$), with the exception of Agreeableness ($p = 0.265$) (Figure S8). The following mean scores were calculated for patients and healthy volunteers, respectively: Openness: $3.6_{\text{patients}}$, $3.6_{\text{healthy volunteers}}$; Conscientiousness: $4.0_{\text{patients}}$, $3.8_{\text{healthy volunteers}}$; Extraversion: $3.5_{\text{patients}}$, $3.5_{\text{healthy volunteers}}$; Agreeableness: $3.5_{\text{patients}}$, $3.3_{\text{healthy volunteers}}$; Neuroticism: $2.6_{\text{patients}}$, $2.5_{\text{healthy volunteers}}$. Patients showed higher Agreeableness than healthy volunteers ($p < 0.05$) and higher age corresponded with higher levels of Conscientiousness. Female sex was directly proportional to Neuroticism ($p < 0.05$) and Conscientiousness ($p < 0.05$). Figures S9–11 summarize correlations between TIPI scores with motives for participation, views on data protection and trust in fields of scientific research. Although some significances were calculated, no significant trend could be detected.
The present study assessed the perception of clinical research among patients and healthy volunteers of clinical trials in a European cohort. While most previous studies have focused on a single aspect of clinical trials in either patients or healthy volunteers, this work provides a comprehensive overview of how both subject groups perceive various areas of clinical research.

Consistent with other literature, the predominant motives for participation among our subjects were altruistic motives and trust in attending doctors [10–14, 19, 20]. Over 80% of all participants strongly agreed or agreed that contribution to scientific research, improvement of medical care for future patients, and great trust in their treating physician were motives for their participation in clinical trials. In addition, the majority (>80%) of all respondents expected researchers, patients, and companies that develop drugs and medicinal products to profit from clinical trials.

According to Good Clinical Practice (GCP) standard, the participation of patients in clinical trials is generally not remunerated. Rather, the opportunity to receive a new treatment alone represents the trial’s benefit and has been documented to be an influential factor for patients [12]. However, our data suggest that health benefits were not primary motivators among the included patients, and fear of disease progression was not of substantial concern. Rather, these motives were outweighed by altruistic considerations and trust in attending physicians. Nevertheless, with age, patients did show increasing interest in receiving the best available
treatment. These findings underscore the importance of the physician–patient relationship during recruitment and the informed consent process, as well as the potential value of highlighting the public benefits derived from the study and the fact that participation in the study will not compromise standard of care when recruiting older patients.

In contrast, it is widely accepted to include monetary incentives in the recruitment of healthy subjects, which are often a necessary component for the successful completion of clinical trials [18, 31]. Furthermore, there is no evidence suggesting that commonly used payments represent undue remuneration or that financial inducements may distort

---

**Fig. 3** Expected beneficiaries of clinical trials. Shown is the response distribution to a question worded as, “How much do you think the following groups would benefit from clinical trials?”

---

**Fig. 4** Concern about the potential consequences of sharing personal information. Shown is the response distribution to a question worded as, “During your participation in clinical trials, you have consented to the disclosure of your personal data to third parties in anonymized form. How concerned are you about the following possible consequences of sharing anonymous clinical trial data?”

© Springer
healthy volunteers’ risk perception associated with the clinical trial [32, 33]. Yet surprisingly, only 56% of healthy volunteers strongly agreed or agreed to have participated due to financial benefits. Thus, although financial compensation is often the focus of recruitment, it may not be the only incentive for subjects to participate in clinical trials. Instead, similarly to patients, the most dominant motives in healthy volunteers were altruistic motives and trust in their attending physicians. Conversely, yet in line with literature, healthy volunteers that indicated financial motives for participation had participated in more clinical trials [34]. This suggests that remuneration may indeed be linked to repeated clinical trial participation in healthy volunteers. Finally, female sex and older age positively correlated with altruistic motives and negatively correlated with financial incentives.

As part of their participation in clinical trials, participants consented to the disclosure of data to third parties in anonymized form. Contrary to public hesitancy [24, 25], subjects did not express considerable concern about the potential risks of data sharing. The perceived risk was lower than that found in previous studies [14, 35]; less than 10% of participants were “very concerned” about any risks mentioned in the questionnaire. The greatest concern was expressed about the misuse of personal data for marketing purposes instead of scientific purposes (8.5% “very concerned”) or for capitalization by companies or individuals (6.4%). A recent systematic review highlights the importance of communication during the process of informed consent and suggests that trust in scientific research and clinicians may mitigate concern regarding data sharing [36]. In agreement with these findings, our subjects expressed considerable trust (76.4% show “great trust” or “trust”) in government research institutions and in medical staff (86.6%), which correlated with lower concern about the consequences of data sharing. Similarly, most participants reported an elaborate, thorough and easily comprehensible process of informed consent. However, we acknowledge that clinical trial participants may represent those, who are least concerned about the potential violations of data protection and most enthusiastic about contributing to scientific progress. Thus, the perception and concerns of data sharing may differ from those of the general public or patients.

When examining personality traits, there was a significant difference between patients and healthy volunteers only in the trait of Agreeableness, with lower scores documented in the healthy volunteers. Although some sporadic significances were calculated when comparing personality traits with motives for participation, views on data sharing and trust in fields or research, no significant trend could be detected. Therefore, the TIPI did not function as an accurate predictor of clinical trial participation behavior. However, this may in part be due the loss of content validity and reliability when using abbreviated measures such as the TIPI [30].

**Limitations**

Most participants took part in phase I clinical trials, thus resulting in a large number of young, healthy individuals. This resulted in an underrepresentation of old (> 60 years of age) participants and an imbalance between healthy volunteers (71.7%) and patients (28.3%). Furthermore, the sample consisted exclusively of individuals that have participated in clinical trials. Therefore, it was not possible to compare our findings to a control group, i.e., subjects that have declined...
participation in the past, or to determine factors responsible for non-participation. Hence, it was also not possible to determine whether the motives that were strongly represented in our sample were decisive for participation in clinical trials or whether these motives were only shared by individuals who generally have an affinity for trial participation. Despite the substantial size of our sample, response rates were low, possibly leading to nonresponse bias. In addition, we acknowledge the bias of selective memory. We were unable to determine the date and time of initial participation; hence, participation may lie far in the past and some data may have been recalled erroneously. Finally, no adjustment was conducted for multiple testing.

Conclusion

In our study, altruistic motives and trust in attending physicians were the predominant motives for clinical trial participation in healthy volunteers and patients. This underscores the importance of the physician–patient relationship during the informed consent process. In addition, with age, patients increasingly expected to receive the best available treatment by participating in trials. Healthy volunteers who reported financial motives for their participation had participated in more clinical trials. Therefore, it may be beneficial to emphasize these benefits during the process of informed consent.

Our subjects expressed little concern about the potential risks of sharing their personal information as part of their participation. Finally, in our study, the Ten-Item Personality Index did not function as an accurate predictor of clinical trial behavior.

Supplementary Information The online version contains supplementary material available at https://doi.org/10.1007/s00228-022-03366-3.

Author contribution F.B., P.M., and M.Z. designed the study. F.B., P.M., L.Y., M.W., T.F., T.W., and M.Z. performed the study. F.B. and L.Y. performed the analysis. F.B. wrote the original draft of the manuscript and created the tables and figures. F.B., P.M., and M.W. managed study documentation and material preparation. All authors contributed to the final version of the manuscript and approved it for publication.

Funding Open access funding provided by Medical University of Vienna.

Data availability The data that support the findings of this study are available from the corresponding author upon reasonable request.

Declarations

Ethics approval Ethical approval for this study was obtained from the Ethics Committee of the Medical University of Vienna. The study was performed in accordance with the ethical standards as laid down in the 1964 Declaration of Helsinki and its later amendments.

Informed consent All participants included in this study underwent a thorough process of informed consent and gave written and oral consent prior to enrollment.

Competing interests The authors declare no competing interests.

References

1. Odgaard-Jensen J, Vist GE, Timmer A, Kunz R, Akli EA, Schünemann H, Briel M, Nordmann AJ, Pregno S, Oxman AD (2011) Randomisation to protect against selection bias in healthcare trials Cochrane Database Syst Rev 4. https://doi.org/10.1002/14651858.MR000012.pub3
2. Eichler HG, Pignattì F, Schwarzer-Daum B et al (2021) Randomized controlled trials versus real world evidence: neither magic nor myth. Clin Pharmacol Ther 109:1212–1218. https://doi.org/10.1002/cpt.2083
3. Collins R, Bowman L, Landray M, Peto R (2020) The magic of randomization versus the myth of real-world evidence. N Engl J Med 382:674–678. https://doi.org/10.1056/NEJMsb1901642
4. Fogel DB (2018) Factors associated with clinical trials that fail and opportunities for improving the likelihood of success: a review. Contemp Clin Trials Commun 11:156–164. https://doi.org/10.1016/j.cctc.2018.08.001
5. Kadam RA, Borde SU, Madas SA et al (2016) Challenges in recruitment and retention of clinical trial subjects. Perspect Clin Res 7:137–143. https://doi.org/10.4103/2229-3485.184820
6. McDonald AM, Knight RC, Campbell MK et al (2006) What influences recruitment to randomised controlled trials? A review of trials funded by two UK funding agencies. Trials 7:9. https://doi.org/10.1186/1745-6215-7-9
7. Newington L, Metcalfe A (2014) Factors influencing recruitment to research: qualitative study of the experiences and perceptions of research teams. BMC Med Res Methodol 14:10. https://doi.org/10.1186/1471-2288-14-10
8. Ford JG, Howerton MW, Lai GY et al (2008) Barriers to recruiting underrepresented populations to cancer clinical trials: a systematic review. Cancer 112:228–242. https://doi.org/10.1002/cncr.23157
9. Treweek S, Lockhart P, Pitkethly M, Cook JA, Kjeldstrøm M, Johansen M, Taskila TK, Sullivan FM, Wilson S, Jackson C, Jones R (2013) Methods to improve recruitment to randomised controlled trials: Cochrane systematic review and meta-analysis. BMJ Open 3. https://doi.org/10.1136/bmjopen-2012-002360
10. Truong TH, Weeks JC, Cook EF, Joffee S (2011) Altruism among participants in cancer clinical trials. Clin Trials 8:616–623. https://doi.org/10.1177/1740774511414444

Springer
11. Jenkins V, Fallowfield L (2000) Reasons for accepting or declining to participate in randomized clinical trials for cancer therapy. Br J Cancer 82:1783–1788. https://doi.org/10.1054/bjoc.2000.1142

12. Godskesen T, Hansson MG, Nygren P et al (2015) Hope for a cure and altruism are the main motives behind participation in phase 3 clinical cancer trials. Eur J Cancer Care 24:133–141. https://doi.org/10.1111/ecc.12184

13. Stunkel L, Grady C (2011) More than the money: a review of the literature examining healthy volunteer motivations. Contemp Clin Trials 32:342–352. https://doi.org/10.1016/j.cct.2010.12.003

14. Mello MM, Lieou V, Goodman SN (2018) Clinical trial participants’ views of the risks and benefits of data sharing. N Engl J Med 378:2202–2211. https://doi.org/10.1056/NEJMsas1713258

15. Martinson BC, Lazovich DA, Lando HA et al (2000) Effectiveness of monetary incentives for recruiting adolescents to an intervention trial to reduce smoking. Prev Med 31:706–713. https://doi.org/10.1006/pmed.2000.0762

16. Jia P, Furuya-Kanamori L, Qin Z-S et al (2020) Association between response rates and monetary incentives in sample study: a systematic review and meta-analysis. Postgrad Med J 97:501-510. https://doi.org/10.1136/postgradmedj-2020-137868

17. Almeida L, Azevedo B, Nunes T et al (2007) Why healthy subjects volunteer for phase I studies and how they perceive their participation? Eur J Clin Pharmacol 63:1085–1094. https://doi.org/10.1007/s00229-007-0368-3

18. Tishler CL, Bartholomae S (2002) The recruitment of normal healthy volunteers: a review of the literature on the use of financial incentives. J Clin Pharmacol 42:365–375. https://doi.org/10.1177/00912700222011409

19. Sood A, Prasad K, Chhatwani L et al (2009) Patients’ attitudes and preferences about participation and recruitment strategies in clinical trials. Mayo Clin Proc 84:243–247. https://doi.org/10.4065/84.3.243

20. Grant CH, Cissna KN, Rosenfeld LB (2000) Patients’ perceptions of physicians communication and outcomes of the accrual to trial process. Health Commun 12:23–39. https://doi.org/10.1207/S15327027HC1201_02

21. Eichler H-G, Pétavy F, Pignatti F, Rasi G (2013) Access to patient-level trial data — a boon to drug developers. N Engl J Med 369:1577–1579. https://doi.org/10.1056/NEJMtp1310771

22. Sampson MG, Kang HM (2019) The path to open data. Nat Rev Nephrol 15:521. https://doi.org/10.1038/s41581-019-0188-6

23. Mello MM, Francker JK, Wilenzick M et al (2013) Preparing for responsible sharing of clinical trial data. N Engl J Med 369:1651–1658. https://doi.org/10.1056/NEJMhle1309073

24. Papoutsi C, Reed JE, Marston C et al (2015) Patient and public views about the security and privacy of electronic health records (EHRs) in the UK: results from a mixed methods study Healthcare Information Systems. BMC Med Inform Decis Mak 15:1–15. https://doi.org/10.1186/s12911-015-0202-2

25. Garrison NA, Sathe NA, Antommaria AH et al (2016) A systematic literature review of individuals’ perspectives on broad consent and data sharing in the United States. Genet Med 18:663–671. https://doi.org/10.1038/gim.2015.138

26. Johnson MO (2000) Personality correlates of HIV vaccine trial participation. Personality Individ Differ 29:459–467. https://doi.org/10.1016/S1091-8869(99)00206-8

27. Goldberg LR (1990) Personality processes and individual differences - an alternative “Description of Personality”: the Big-Five factor structure. J Pers Soc Psychol 59:1216–1229. https://doi.org/10.1037/0022-3514.59.6.1216

28. Halpern SD, Rentfrow PJ, Swann WB (2003) A very brief measure of the Big Five personality domains. J Res Pers 37:504–528. https://doi.org/10.1016/S0022-4825(03)00046-1

29. Crédé M, Harms P, Niehorster S, Gaye-Valentine A (2012) An evaluation of the consequences of using short measures of the Big Five personality traits. J Pers Soc Psychol 102:874–888. https://doi.org/10.1037/a0027403

30. Grady C (2005) Payment of clinical research subjects. J Clin Invest 115:1681–1687. https://doi.org/10.1172/JCI25694

31. Bentley JP, Thacker PG (2004) The influence of risk and monetary payment on the research participation decision making process. J Med Ethics 30:293–298. https://doi.org/10.1136/jme.2002.001594

32. Halpern SD, Karlawish JHT, Casarett D et al (2004) Empirical assessment of whether moderate payments are undue or unjust inducements for participation in clinical trials. Arch Intern Med 164:801–803. https://doi.org/10.1001/archinte.164.7.801

33. Fisher JA, McManus L, Wood MM et al (2018) Healthy volunteers’ perceptions of the benefits of their participation in phase I clinical trials. J Empir Res Hum Res Ethics 13:494–510. https://doi.org/10.1177/1556264618804962

34. Whidett R, Hunter I, Engelbrecht J, Handy J (2006) Patients’ attitudes towards sharing their health information. Int J Med Informatics 75:530–541. https://doi.org/10.1016/j.ijmedinf.2005.08.009

35. Hutchesons EL, Loomes M, Butow P, Boyle FM (2021) A systematic literature review of attitudes towards secondary use and sharing of health administrative and clinical trial data: a focus on consent. Syst Rev 10:1–41. https://doi.org/10.1186/s13643-021-01663-z

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