Religious Perspectives on Precision Medicine in Singapore

Hui Jin Toh1 · Angela Ballantyne1,2 · Serene Ai Kiang Ong1 · Chitra Sankaran3 · Hung Yong Tay4 · Malminderjit Singh5 · Raza Zaidi6 · Roland Chia7 · Sarabjeet Singh8 · Swami Samachittananda9 · You Guang Shi10 · Zhixia Tan4 · Tamra Lysaght1

Received: 6 July 2021 / Revised: 6 July 2021 / Accepted: 7 July 2021 / Published online: 6 September 2021 © The Author(s) 2021

Abstract

Precision medicine (PM) aims to revolutionise healthcare, but little is known about the role religion and spirituality might play in the ethical discourse about PM. This Perspective reports the outcomes of a knowledge exchange fora with religious authorities in Singapore about data sharing for PM. While the exchange did not identify any foundational religious objections to PM, ethical concerns were raised about the possibility for private industry to profiteer from social resources and the potential for genetic discrimination by private health insurers. According to religious authorities in Singapore, sharing PM data with private industry will require a clear public benefit and robust data governance that incorporates principles of transparency, accountability and oversight.

Keywords  Precision medicine · Religious views · Health data sharing · Genomics · Private sector · Health equity

* Hui Jin Toh
medthj@nus.edu.sg

1 Centre for Biomedical Ethics, Yong Loo Lin School of Medicine, National University of Singapore, Singapore
2 Department of Primary Health Care and General Practice, University of Otago, Wellington, New Zealand
3 Hindu Centre, Singapore
4 Taoist Mission (Singapore), Singapore
5 Sikh Advisory Board, Singapore
6 Jaafari Muslim Association Singapore, Singapore
7 National Council of Churches of Singapore, Singapore
8 Young Sikh Association (Singapore), Singapore
9 Ramakrishna Mission, Singapore
10 Singapore Buddhist Federation, Singapore
Background

Governments around the world are investing in large scale national programmes for precision medicine (PM) and Singapore has recently launched the National Precision Medicine (NPM) strategy (PRECISE n.d.). While definitions can vary, PM essentially aims to optimise health outcomes by using knowledge about genomics, lifestyle, and environment factors to better tailor medicine and healthcare (Wang et al. 2016). Unlike the traditional medical practice of prescribing similar treatment plans for patients with common clinical presentations, PM stratifies patients into sub-groups thereby facilitating more precise diagnoses and targeted therapies.

Public attitudes are important for gauging the social licence needed to conduct data-intensive research in the absence of specific consent. PM requires broad public buy-in and cooperation. To understand how genomics, lifestyle, and environment factors interact, researchers need access to substantial amounts of data containing information about clinical observations, genomics and health outcomes from a diverse range of volunteers. Tracking the health of individuals and groups over time requires sophisticated infrastructure to collect, store, link, and re-purpose data for analysis. Given the scope of PM research, multiple parties in both the private and public sectors are typically involved, including public health services, academic researchers, private sector pharmaceutical and biotechnology companies, private insurance companies, and technology companies (Wang et al. 2016). Longitudinal large-scale data collection and sharing across the public and private sector raise several challenges including data security, privacy, public trust, as well as the scope of data-subjects’ consent and social licence for data sharing (Chataway et al. 2012).

This Perspective summarises the outcomes of knowledge exchange in a facilitated workshop with religious authorities in Singapore (see Annex A in the Supplementary Material) to share their perspectives on PM, and data sharing with the organizations in the public and private sectors. To date, there has been little published about the role of religion and spirituality in relation to the ethical challenges of PM (Prosperi et al. 2018; Fisher et al. 2020) or public attitudes to PM (Yeary et al. 2020; Sanderson et al. 2017). Broadly speaking, we know that religious views and practices influence health behaviours and treatment efficacy (Fisher et al. 2020; Ogden 2016) and that religiosity has been associated with a lack of faith in science (Johnson et al. 2015). Sanderson et al. (2017) found that “very religious” respondents in the USA were less willing to participate in biobanking research than non-religious participants; and therefore recommend that PM programmes actively address the concerns of religious communities to ensure diverse representation in PM cohorts. Sheppard et al. (2018) also found that religiosity was associated with lower rates of biospecimen donation for PM-related research.

It is worth noting that this prior research on religion and PM (Yeary et al. 2020; Sanderson et al. 2017; Sheppard et al. 2018), or more broadly on religiosity and views on genetic testing and genomic research (Kinney et al. 2006; Lee et al. 2019), are nearly exclusively based in the USA and may not be reflective of the relationship (if any) between religiosity and views on PM in Singapore or other Asian countries. Therefore, it is important to consider whether PM research and governance
processes need to address specific religious considerations in relation to PM in Singapore. Research shows that religious leaders and proscriptions can be influential in their congregations’ behaviours and decisions about health interventions (Thomas et al. 2015; Hungerman 2012). A study in Jordan found that religious permission for biobanking\(^1\) was one of the two most influential factors for prompting public participation in biobanking (Ahram et al. 2014). For these reasons, we invited religious authorities from across Singapore to share knowledge in a workshop on ethical issues raised by PM. Outcomes from this workshop will be of interest to academics and policymakers responsible for establishing the social licence for PM.

**A Workshop with Religious Authorities in Singapore**

This Perspective presents the main outcomes of the workshop. Many of the ethical and governance issues raised during the discussions correspond with prior research on public attitudes towards PM globally (Holden et al. 2019). Here, four broad themes are highlighted: (i) religious values and support for PM in foundational principles, scripture or teaching; (ii) potential religious concerns/challenges for PM; (iii) potential concerns associated with sharing PM data with commercial or private partners; and (iv) recommendations for governance approaches in PM in Singapore. There was no outright disagreement between religious leaders on any points discussed; rather representatives of particular religions choose to speak to certain points, or emphasise particular values or religious tenets.

**Religious Values and Support for PM in Foundational Principles, Scripture or Teaching**

The discussions suggested no foundational religious objections to PM or scripture that would directly prevent participation in PM. Rather, the representatives in attendance emphasized values of charity, community, assisting others and acting for the greater good, which were perceived to lend support for participation in PM research. Participating in PM research could be viewed as a way of fulfilling a duty to help those sick or suffering.

*securing public benefit and safeguarding humanity’s well-being, is of utmost importance.* (Muslim representative)

However, equity and access to medicine, specifically the products of PM, remain significant considerations. The increasing cost of medicine is already a concern in Singapore (How and Fock 2014), and many current medical treatments and procedures are widely viewed to be out of the reach of average Singaporeans. This

---

\(^1\) Some representatives were shown a statement indicating that religion does not prohibit individuals from donating biospecimens for research and biobanking.
disparity creates inequality between individuals, which is likely to be at odds with some religious teachings. Taoism, for example, is aligned with relational justice through its emphasis on justice as “dao” (way-making) (Tan 2019). This implies that all people are of equal worth and, therefore, should have equal access to medicine that potentially helps to save lives.

*all these high cutting-edge medicine which costs is not something which is affordable for most people. In a way, you definitely do [have] discrimination between the haves and the have nots.* (Taoist representative)

### Potential Religious Concerns/Challenges Regarding PM

There was general consensus that people contributing data to PM ought to undergo a robust informed consent process, and that PM data should not be used or shared without consent. Minimally, broad consent is needed at the point of recruitment into the PM programme. Broad consent is an alternative to specific consent and focuses on describing the general purpose and scope of the research programme. Broad consent should include sufficient information to enable a reasonable person to reach a decision about whether to consent or decline to participate; but this need not involve specifying every conceivable research use (Maloy and Bass 2020). These broad consent approaches are typically used in PM because of the difficulty in describing all possible future uses of data at the point of recruitment into a PM programme and the high cost (in terms of time, human resources and funding) of concurrently seeking consent for each data use (Schaefer et al. 2019). Under the Singaporean Human Biomedical Research Act 2015, broad consent can be obtained in lieu of specific consent for storage and secondary uses of identifiable data and biospecimens (Goh 2018). It is becoming generally accepted that genomic data cannot be truly anonymised (Shabani and Marelli 2019). The consent process for participants entering a PM programme of research must therefore include information about who the data may be shared with, the process of de-identification, and the potential risk of re-identification, even where this risk is small.

More specifically, the representatives from Catholicism and Hinduism emphasized rationality, free will, autonomy and personal responsibility. The following Catholic teaching demonstrating the relationship between morality, responsibility and voluntariness: “Freedom characterizes properly human acts. It makes the human being responsible for acts of which he is the voluntary agent. His deliberate acts properly belong to him” (Catholic Church 2000, 1745). Hinduism also places importance on the individual and individual will. For example, Hindu rituals and practises are diverse with significant freedom for individual choice and preference (Harman 2004); and Hindu ethical teaching focuses on the “cultivation of the self” (Baird 1998) towards the goal of non-attachment.

*as far as Hinduism is concerned, it gives a lot of importance to the individual and individual will* (Hindu representative)
From these perspectives, the absence of specific consent to each instance of data sharing in a PM programme could be seen as a potential barrier to individuals’ ability to make decisions and responsibly exercise their free will. The value of individual autonomy and responsibility in Catholicism and Hinduism is in prima facie conflict with broad-, blanket- and future-consent models in PM. Yet, representatives recognized that medical technology and genomic research methods develop; therefore, not all future uses of PM data could reasonably be predicted at the time of enrolment in a PM programme. Some suggested that, in order to respect individual autonomy, periodic re-consent may be preferable to relying solely on broad consent at the time of enrolling in the programme. During the re-consenting process, significant changes in the scope of sharing, linking and new uses of the data should be communicated to participants in a PM programme.

Representatives suggested that using data outside the parameters of the individual’s consent would raise moral challenges. In the absence of specific consent, transparency and the right to withdraw from a PM programme were seen as important mechanisms for promoting autonomy and free will as embodied in religious teachings. The ethical need for transparency is closely linked to the value of human autonomy, which allows individuals the opportunity to make decisions independently by weighing information in light of their convictions and values (Felzmann et al. 2019). Transparency may help ensure that participants can access knowledge about how their data is being used and enable them to withdraw from the programme if they believe that data uses are outside the parameters of their original consent or their current expectations.

**Potential Concerns Associated with Sharing Data With Commercial or Private Partners**

Representatives at the workshop expressed scepticism towards data sharing with commercial or private partners, including pharmaceutical and biotechnology companies, private insurance companies, and technology companies. As indicated above, charity and community benefit were central pillars for all representatives present at the workshop. While this focus on charity provides impetus to participate in a PM programme that aims to improve community health and wellbeing, they explained that it also underpins a strong reluctance to allow the commercialisation and commodification of PM data, and private profiteering from social resources. Commercialisation and commodification of PM data can be interpreted as treating PM data as “an asset” and selling or distributing PM data for private financial gains (Bottis and Bouchagiar 2018).
if it’s for commercial use, and for profiteering from the information, that we do not support also. But if it’s for something that medicine to cure that will help humankind, we are for it. (Taoist representative)

the most important part is to assure that the personal information or the information obtained will not be used for any other purpose other than people consented for or for commercial gains. While the intent of benefitting humanity is commendable, it needs to be ensured that the use of personal data for purposes other than service to humanity will not happen. (Muslim representative)

International empirical research has consistently demonstrated that the public and patients are reluctant to allow the use of health-related data for commercial purposes (Shabani et al. 2014; Hill et al. 2013; Kalkman et al. 2019; Grant et al. 2013). The NICE Citizens Council (2015) in the UK found that half of the Council had concerns about anonymous data\(^2\) being shared widely, and these related in part to objections to data being sold on to other organisations, used for profit, or for commercial purposes other than research. In response, some data access policies limit access by commercial entities (Shabani and Borry 2016). There was concern expressed at the workshop that sharing PM data with commercial companies could also facilitate genetic discrimination by insurance companies, and inequity arising from the development of expensive and inaccessible drugs by pharmaceutical companies (Lee et al. 2019).

From a Catholic perspective, health may be viewed as a fundamental right. Pope Benedict XVI has been emphatic on the need to “guarantee adequate [health] care to all” (Sibley 2016). As such, health should not be left to the market to determine or distribute.

**Health is a fundamental right. It’s not like they’re offering other kinds of things that are optional for people.** (Catholic representative)

This reasoning shares commonalities with the work of bioethicists Norman Daniels (1985) who has written about the role of health as a primary good. Daniels uses the Rawlsian (Rawls 1999) framework of primary goods and fair equality of opportunity to draw attention to the specific importance of health care. Daniels argues that health care deserves special attention because of the unequal natural distribution of health care needs and because of the strategic importance of health to an individual’s opportunity range and future life choices. On this view, justice requires that institutions ensure that the social basis of health should be distributed fairly within the community. This means that access to the benefits of social institutions that affect people’s ability to protect and maintain their health, arguably including PM programmes, should be fairly distributed amongst the community. It was reasoned at the workshop that for-profit companies and commercial interests in PM should not be permitted to distort or unfairly determine access to health resources.

---

\(^2\) There is growing consensus that true anonymity of data is now technically impossible; therefore, we prefer to talk about degrees of data de-identification and security rather than anonymous data.
Nonetheless, there was some openness to Public Private Partnerships (PPPs) or data-sharing arrangements that produce public benefit in addition to private profit. For example, Catholicism accepts the role of private property, commercial interests and economic role of markets, so long as these are arranged to serve the higher purposes of allowing individuals to express their free will and aligned with the common good and communal purpose (Koterski 2012).

there is nothing wrong with profit. [If] the profit is done by a private company for the common good, they deserve that profit. (Catholic representative)

PPPs can be an effective method of harnessing the potential of health and genomic data and can include public and private sector partners working across the data chain—producing, analysing, using or repurposing data (Ballantyne and Cameron 2019). Programmes such as the Innovative Medicines Initiative’s (IMI) established by the European Union works as an international data sharing platform aiming to stimulate novel drug development. Some of the most successful biobanks, such as UK Biobank (n.d.) and FinnGen (n.d.), allow commercial and private sector organisations access to their data. Religious representatives involved in this workshop expressed openness to public private partnerships or data-sharing arrangements with commercial companies that produce public benefit in addition to private profit.

**Recommendations for Governance Approaches in PM**

The general consensus at the workshop was that a commitment to public benefit, transparency, accountability and robust governance models will be necessary to engender the support of religious authorities for the sharing of PM data from Singapore with commercial companies, especially in the absence of specific consent for such sharing. Trustworthy governance should include a review process for data sharing arrangements to ensure the public benefit requirement is met. This review process should include lay perspectives. Existing oversight and approval mechanisms for health data sharing include Data Access Committees (DACs) and Institutional Review Boards (IRB). However, in Singapore, IRB approval would not be required for sharing de-identified/non-personal PM data with research partners, including private industry. The Personal Data Protection Act (PDPA) and the Human Biomedical Research Act (HBRA) apply only to personal (reasonably identifiable) data sharing. Workshop discussions concluded that, given the national interests in protecting and stewarding Singaporean genomic data, establishing a national PM oversight body or panel responsible for making decisions about when to share PM data, particularly if private industry is involved, may help to engender trust within religious communities about data sharing arrangements.

---

3 DACs are a group that is charged with the responsibility of reviewing and assessing data access requests; they have a dual responsibility with respect to “both promotion of data sharing and protection of data subjects, their communities, data producers, their institutions and the scientific enterprise” (Cheah and Piasecki 2020). IRBs (in some jurisdictions these are called Research Ethics Committees) are charged with reviewing and approving human subjects research.
There was also consensus about data sharing being guided by the principles of openness, transparency, honesty and accountability. These norms were seen as essential pillars for a trustworthy and religiously acceptable PM programme. Openness — with respect to ownership of the data, users of the data, place of storage, and rights retained by the donors of the data — would demonstrate respect to those who contribute to PM:

*As far as the Christian faith is concerned, it does come back to the dignity of the human being who has contributed those data.* (Christian representative)

To further facilitate transparency and accountability, an audit trail for the data sharing should be publicly available. The outcomes of discussions suggested that this openness includes a list of all PM projects, including collaborations or public–private partnerships using the data. The UK Biobank (n.d.) takes this approach: projects using the data are described on their website. Obligations of transparency may conflict with commercial confidentiality with respect to some public–private partnerships; nonetheless representatives stressed that transparency is a high priority for ensuring moral legitimacy and public trust. The data governance principles raised throughout the conversations, included accountability, transparency, and dignity/respect, align with those described elsewhere (Merson et al. 2015; Xafis et al. 2020) and will be an important component of data governance structures for PM.

**Conclusions**

This Perspective describes the outcomes of a workshop for knowledge exchange between representatives of the major religions in Singapore on PM and sharing data with public agencies and commercial companies. The general consensus to emerge from the workshop was that PM could be an important contributor to the common good that should be embedded with values of charity, community, respect and trust. Securing the social license would thus require robust and trustworthy systems of governance and oversight to ensure transparency, accountability and the equitable distributions of benefits. This consensus accords with the bioethical principles and norms identified in the prior theoretical and empirical literature on health data governance.

**Supplementary Information** The online version contains supplementary material available at https://doi.org/10.1007/s41649-021-00180-4.

**Acknowledgements** We thank the other participants who took part in the workshop but are not credited with authorship. We also thank Professor Tai E Shyong from National University Hospital who was present during the workshop to address technical questions from the participants.

**Funding** This research is supported by the Ministry of Education, Singapore, under its Social Science Research Council Thematic Grant (MOE2017-SSRTG-028).

**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

Ahram, Mamoun, Areej Othman, Manal Shahrouri, and Mustafa Ebtihal. 2014. Factors influencing public participation in biobanking. European Journal of Human Genetics 22: 445–451. https://doi.org/10.1038/ejhg.2013.174.

Baird, Robert D. 1998. Hinduism. In Encyclopedia of Applied Ethics, ed. Ruth Chadwick, 582–592. Oxford: Elsevier.

Ballantyne, Angela and Cameron Steward. 2019. Big data and public-private partnerships in healthcare research: the application of an ethics framework for big data in health and research. Asian Bioethics Review 11(3): 315–326. https://doi.org/10.1007/s41649-019-00100-7.

Bottis, Maria, and George Bouchagiar. 2018. Personal data v. big data: Challenges of commodification of personal data. Open Journal of Philosophy 8: 206–215. https://doi.org/10.4236/ojpp.2018.83015.

Catholic Church. 2000. Catechism of the Catholic Church. 2nd ed. Vatican City: Libreria Editrice Vaticana.

Chataway, Joanna, Caroline Fry, Sonja Marjanovic, and Ohid Yaqub. 2012. Public-private collaborations and partnerships in stratified medicine: Making sense of new interactions. New Biotechnology 29 (6): 732–740. https://doi.org/10.1016/j.nbt.2012.03.006.

Cheah, Phaik Yeong, and Jan Piasecki. 2020. Data Access Committees. BMC Medical Ethics 21: 12. https://doi.org/10.1186/s12910-020-0453-z.

Daniels, Norman. 1985. Just Health Care. Cambridge: Cambridge University Press.

Felzmann, Heike, Fosch Villaronga Eduard, Christoph Lutz, and Aurelia Tamò-Larrieux. 2019. Robots and transparency: The multiple dimensions of transparency in the context of robot technologies. IEEE Robotics & Automation Magazine 26 (2): 71–78. https://doi.org/10.1109/MRA.2019.2904644.

FinnGen. n.d. https://www.finngen.fi/en. Accessed 3 Feb 2021.

Fisher, Elena R., Rebekah Pratt, Riley Esch, Megan Kocher, Katie Wilson, Whiwon Lee, and Heather A Zierhut. 2020. The role of race and ethnicity in views toward and participation in genetic studies and precision medicine research: A systematic review of qualitative and quantitative studies. Molecular Genetics & Genomic Medicine 8 (2): e1099. https://doi.org/10.1002/mgge.31099.

Goh, Yeow Tee. 2018. The Human Biomedical Research Act: Compliance and Risks. https://centres.sg/wp-content/uploads/2017/10/PO3_AProf-YT-GOH-release-01_02-Mar-18_13Mar18.pdf. Accessed 28 Apr 2021.

Grant, Aileen, Jenny Ure, Donald Nicolson, Janet Hanley, Aziz Sheikh, Brian McKinsity, et al. 2013. Acceptability and perceived barriers and facilitators to creating a national research register to enable ‘direct to patient’ enrolment into research: The Scottish Health Research Register (SHARE). BMC Health Services Research 13 (1): 422. https://doi.org/10.1186/1472-6963-13-422.

Harman, William. 2004. Hindu Devotion. In Contemporary Hinduism: Ritual, Culture, and Practice, ed. Robin Rinehart, 99–122. Santa Barbara, CA: ABC-CLIO.

Hill, Elizabeth M., Emma L. Turner, Richard M. Martin, and Jenny L. Donovan. 2013. “Let’s get the best quality research we can”: Public awareness and acceptance of consent to use existing data in health research: A systematic review and qualitative study. BMC Medical Research Methodology 13 (1): 72. https://doi.org/10.1186/1471-2288-13-72.

Holden, Ciara, Lauren Bignell, Somnath Mukhopadhyay, and Christina Jones. 2019. The public perception of the facilitators and barriers to implementing personalized medicine: A systematic review. Personalised Medicine. https://doi.org/10.2217/pme-2018-0151.

How, Choon How, and Kwong Ming Fock. 2014. Healthcare in Singapore: the present and future. Singapore Medical Journal 55 (3): 126–127. https://doi.org/10.11622/smedj.2014027.

Hungerman, Daniel M. 2012. Do religious proscriptions matter? Evidence from a theory-based test. Journal of Human Resources 49 (4): 1053–1093. https://doi.org/10.3368/jhr.49.4.1053.
Johnson, David R., Christopher P. Scheitl, and Elaine Howard Ecklund. 2015. Individual religiosity and orientation towards science: reformulating relationships. *Sociological Science* 2: 106–124. https://doi.org/10.15195/v2.a7.

Kalkman, Shona, Johannes van Delden, Banerjee Amitava, Benoît Tyl, Menno Mostert, and Ghislaine van Thiel. 2019. Patients’ and public views and attitudes towards the sharing of health data for research: A narrative review of the empirical evidence. *Journal of Medical Ethics*. https://doi.org/10.1136/medethics-2019-105651.

Kinney, Anita Yeomans, Sara Ellis Simonsen, Bonnie Jeanie Baty, Diptasri Mandal, Susan L. Neuhausen, Kate Seggar, Rich Holubkov, et al. 2006. Acceptance of genetic testing for hereditary breast ovarian cancer among study enrollees from an African American kindred. *American Journal of Medical Genetics Part A* 140A: 813–826. https://doi.org/10.1002/ajmg.a.31162.

Koterski, Joseph. 2012. Human nature from a Catholic perspective. *American Journal of Economics and Sociology* 71 (4): 809–39. https://doi.org/10.1111/j.1536-7150.2012.00840.x.

Lee, Sandra S-J., Mildred K. Cho, Stephanie A. Kraft, Nina Varsava, Katie Gillespie, Kelly E. Ormond, Benjamin S. Wilfond, et al. 2019. “I don’t want to be Henrietta Lacks”: Diverse patient perspectives on donating biospecimens for precision medicine research. *Genetics in Medicine* 21 (1): 107–113. https://doi.org/10.1038/s41436-018-0032-6.

Merson, Laura, Viet Phong Tran, Nguyen Thanh Nhan Le, Thanh Dung Nguyen, Thi Dieu Ngan Ta, Van Kinh Nguyen, Michael Parker and Susan Bull. 2015. Trust, respect, and reciprocity: informing culturally appropriate data-sharing practice in Vietnam. *Journal of Empirical Research on Human Research Ethics* 10 (3): 251–63. https://doi.org/10.1177/1556266415592387.

Maloy, John W., and Pat F. Bass 3rd. 2020. Understanding broad consent. *Ochsner Journal* 20 (1): 81–86. https://doi.org/10.31486/oj.19.0088.

NICE Citizens Council. 2015. What ethical and practical issues need to be considered in the use of anonymised information derived from personal care records as part of the evaluation of treatments and delivery of care? NICE citizens Council reports No. 18. London: National Institute for Health and Care Excellence (NICE).

Ogden, Joy. 2016. Religious constraints on prescribing medication. *Prescriber* 27 (12): 47–51. https://doi.org/10.1002/psb.1524.

PRECISE. n.d. Precision Health Research Singapore. https://www.npm.sg/about-us/our-story/. Accessed 3 Feb 2021.

Prosperi, Mattia, Jae S. Min, Jiyan Bian, and Francois Modave. 2018. Big data hurdles in precision medicine and precision public health. *BMC Medical Informatics and Decision Making* 18: 139. https://doi.org/10.1186/s12911-018-0719-2.

Rawls, John. 1999. *A Theory of Justice*. Revised. Cambridge, MA: Harvard University Press.

Sanderson, Saskia C., Kyle B. Brothers, Nathaniel D. Mercaido, Ellen Wright Clayton, Armand H. Antommaria, Aufox Matheny, A. Sharon, et al. 2017. Public attitudes toward consent and data sharing in biobank research: A large multi-site experimental survey in the US. *American Journal of Human Genetics* 100: 414–427. https://doi.org/10.1016/j.ajhg.2017.01.021.

Schaefer, G. Owen, E Shyong Tai, and Shirley Sun. 2019. Precision medicine and big data. *Asian Bioethics Review* 11 (3): 275–288. https://doi.org/10.1007/s41649-019-00094-2.

Shabani, Mahsa, Louise Bezuidenhout, and Pascal Borry. 2014. Attitudes of research participants and the general public towards genomic data sharing: A systematic literature review. *Expert Review of Molecular Diagnostics* 14 (8): 1053–1065. https://doi.org/10.1586/14737159.2014.961917.

Shabani, Mahsa, and Luca Marelli. 2019. Re-identifiability of genomic data and the GDPR. *MBO Reports* 20 (6): e48316. https://doi.org/10.15252/embr.201948316.

Shabani, Mahsa, and Pascal Borry. 2016. “You want the right amount of oversight”: Interviews with data access committee members and experts on genomic data access. *Genetics in Medicine* 18 (9): 892–897. https://doi.org/10.1038/gim.2015.189.

Sheppard, Vanessa B., Alejandra Hurtado-de-Mendoza, Yun-Ling Zheng, Ying Wang, Kristi D. Graves, Tania Lobo, Hanfei Xu, et al. 2018. Biospecimen donation among black and white breast cancer survivors: Opportunities to promote precision medicine. *Journal of Cancer Survivorship* 12: 74–81. https://doi.org/10.1007/s11764-017-0646-8.

Sibley, Augus. 2016. Health care’s ills: A Catholic diagnosis. *Linacre Quarterly* 83 (4): 402–422. https://doi.org/10.1080/00243639.2016.1249264.

Tan, Charlene. 2019. Conceptualising social justice in education: a Daoist perspective. *Compare: A Journal of Comparative and International Education* 51 (4): 596–611. https://doi.org/10.1080/0305925.2019.1660144.
Thomas, Tami, Blumling Amy, and Augustina Delaney. 2015. The influence of religiosity and spirituality on rural parents’ health decision making and human papillomavirus vaccine choices. *Advances in Nursing Science* 38 (4): E1–E12. https://doi.org/10.1097/ans.0000000000000094.

UK Biobank. n.d. Approved Research. https://www.ukbiobank.ac.uk/enable-your-research/approved-research. Accessed 3 Feb 2021.

Wang, Zheng-Guo, Liang Zhang, and Wen-Jun Zhao. 2016. Definition and application of precision medicine. *Chinese Journal of Traumatology* 19 (5): 249–250. https://doi.org/10.1016/j.cjtee.2016.04.005.

Xafis, Vicki, G. Owen Schaefer, Markus Labude, Yujia Zhu, and Li Yan Hsu. 2020. The perfect moral storm: Diverse ethical considerations in the COVID-19 Pandemic. *Asian Bioethics Review* 12 (2): 65–83. https://doi.org/10.1007/s41649-020-00125-3.

Yeary, Karen H.K., Kassandra I. Alcaraz, Kimlim Tam Ashing, Chungyi Chiu, Shannon M. Christy, Katarina Friberg Felsted, Qian Lu, et al. 2020. Considering religion and spirituality in precision medicine. *Translational Behavioural Medicine* 10 (1): 195–203. https://doi.org/10.1093/tbm/ibz105.

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