

Expert Insights

—

# Exploring quantum computing use cases for life sciences

Decoding secrets  
of genomes, drugs,  
and proteins

IBM Institute for  
Business Value



## Experts on this topic



### **Dr. Frederik Flöther**

Life Sciences and Healthcare Lead  
IBM Quantum Industry Consulting  
IBM Services  
[linkedin.com/in/frederikfloether](https://www.linkedin.com/in/frederikfloether)  
[frederik.floether@ch.ibm.com](mailto:frederik.floether@ch.ibm.com)

Frederik Flöther is an IBM Quantum Industry Consultant and globally leads efforts for the Life Sciences and Healthcare sector. Frederik is an IBM Academy of Technology member and Senior Inventor. He has deep expertise in quantum computing and artificial intelligence and works with clients to create value through these next-generation technologies. Frederik has authored more than 20 patents, peer-reviewed publications, and white papers.



### **Christopher Moose**

Partner, Life Sciences  
and Healthcare  
IBM Services  
[linkedin.com/in/chrismoose](https://www.linkedin.com/in/chrismoose)  
[chris.moose@us.ibm.com](mailto:chris.moose@us.ibm.com)

Christopher Moose is a Partner in IBM's Healthcare and Life Sciences practice. He has 20 years of global experience advising companies and supporting their operations in more than 40 countries. He also has first-hand care delivery experience by chairing the board of a nonprofit child welfare organization.



### **Ivano Tavernelli**

Global Leader, Advanced Algorithms  
for Quantum Simulations  
IBM Research  
[linkedin.com/in/tavernelli-ivano-1b788086](https://www.linkedin.com/in/tavernelli-ivano-1b788086)  
[ITA@zurich.ibm.com](mailto:ITA@zurich.ibm.com)

Ivano Tavernelli is a Research Staff Member in the Cognitive Computing and Industry Solutions department at IBM Research—Zurich since 2014. In 2018 he became IBM Global leader for Advanced Algorithms for Quantum Simulations, responsible for quantum simulations and applications in chemistry, physics, material science, and optimization problems. He also leads the development of the IBM software platform Qiskit Aqua Chemistry and Physics. His focus is the design of efficient quantum algorithms for state-of-the-art quantum devices based on superconducting circuits.

The authors thank Dr. Panagiotis Barkoutsos, Dr. Wendy Cornell, and Dr. Kayla Lee for their contributions in developing this report.

Quantum computing isn't merely about speed. It's about tackling problems differently and making the seemingly impossible possible, if not commonplace.

## Key takeaways

### **Making the impossible possible**

Quantum computing will be used extensively by new categories of professionals and developers to solve problems once considered unsolvable.

### **Disruptive life sciences use cases**

In the life sciences industry, quantum computing is expected to enable a range of disruptive use cases by linking genomes with outcomes, enhancing drug discovery, and improving protein folding predictions.

### **The time to act is now**

Life sciences could benefit significantly from quantum computing. However, much of the early intellectual property in quantum computing may be proprietary, raising the urgency to get started and engage with partners and ecosystems.

—

## Solving the unsolvable

No doubt you've heard it before. Quantum computers can do some things more efficiently than classical computers. Is that a big deal? Why are advanced computational approaches even needed?

In life sciences, major challenges include understanding the relationships among sequence, structure, and function and how biopolymers interact with one another as well as with small organic molecules that are native to the body or designed as drugs. Such problems are computationally complex and are at the heart of genomic analysis, drug design, and protein folding predictions.

Consider drug design. The number of molecules made up of say 50 atoms that can be built using just 10 different types of atoms amounts to around  $10^{50}$ .<sup>1</sup> If we also factor in the number of possible molecular configurations and conformations that can be sampled at room temperature, the total number of molecules that could potentially constitute a valid drug is exponentially greater than the roughly  $10^{80}$  atoms in the observable universe. Tackling this level of complexity is far beyond the capabilities of classical computers; however, quantum computers could make inroads.

The famous physicist, Richard Feynman, suggested back in the 1980s that "if you want to make a simulation of nature, you'd better make it quantum mechanical."<sup>2</sup> It's about tackling problems differently and making the seemingly impossible possible, if not commonplace.



## Insight: Bits and qubits

Quantum computers process information in a fundamentally different way from traditional computers. Previous computer technology advancements—such as integrated circuits—enabled faster computing but were still based on classical information processing. Quantum computers manipulate quantum bits (qubits). These are unlike classical bits, which store information as either a 0 or 1, as they can display uniquely quantum properties, such as entanglement. As a result, it becomes possible to construct quantum algorithms that can outperform their classical counterparts, since they are not able to leverage quantum phenomena.

Quantum computers could be particularly useful in tackling problems that involve:

- Chemistry, machine learning/artificial intelligence, optimization, or simulation tasks. In fact, machine learning has shown potential to be enhanced by quantum computing and is already helping drive quantum advances<sup>4</sup>
- Complex correlations and interdependencies among many highly interconnected elements, such as molecular structures in which many electrons interact
- Inherent scaling limits of relevant classical algorithms. For instance, the resource requirements of classical algorithms may increase exponentially with problem size, as is the case when simulating the time evolution of quantum systems.<sup>5</sup>

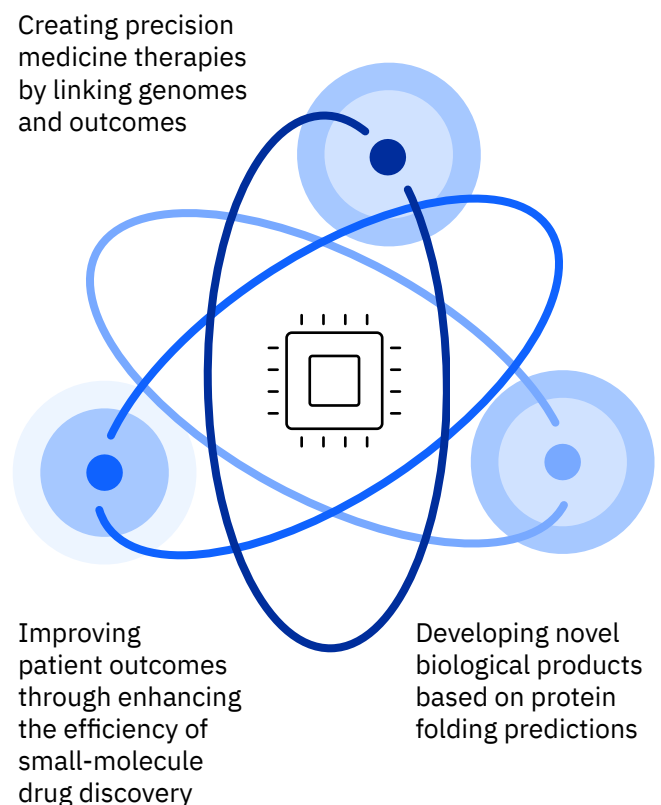
As a result, there is now a cross-industry race toward quantum applications. Within five years, it is possible quantum computing will be used extensively by new categories of professionals and developers to solve problems once considered unsolvable.<sup>3</sup> In the life sciences industry, quantum computing is expected to enable a range of disruptive use cases. These include:

1. Creating precision medicine therapies by linking genomes and outcomes
2. Improving patient outcomes through enhancing the efficiency of small-molecule drug discovery
3. Developing novel biological products based on protein folding predictions (see Figure 1).

—

### Figure 1

Quantum computers may enable three key life sciences use cases that reinforce each other in a virtuous cycle



Research focus has shifted to taking advantage of new computational tools to deepen our understanding of how genomic sequences translate to function.

**Use Case 1: Creating precision medicine therapies by linking genomes and outcomes**

The 15-year, USD 2.7 billion investment to accurately sequence the human genome and subsequent reductions in sequencing costs helped launch the “-omics” era.<sup>6</sup> Accordingly, understanding primary sequences is no longer a major limitation for scientists. Instead, research focus has shifted to taking advantage of new computational tools to deepen our understanding of how genomic sequences translate to function. However, this task is extremely difficult with traditional methods due to the size of the human genome (about 3 billion DNA base pairs), the variation that exists across populations, and the wide range of health outcomes.<sup>7</sup>

Potential opportunities at the intersection of genomics and quantum computing include:<sup>8</sup>

- *Motif discovery and prediction*<sup>9</sup>: DNA, RNA, and amino acid sequences have all been shaped through evolutionary pressures. One bioinformatic challenge is identifying motifs in these sequences, such as patterns that activate or inhibit gene expression and, thereby, help us better understand mechanisms of gene regulation. Classical algorithms to identify motifs are computationally expensive because they require

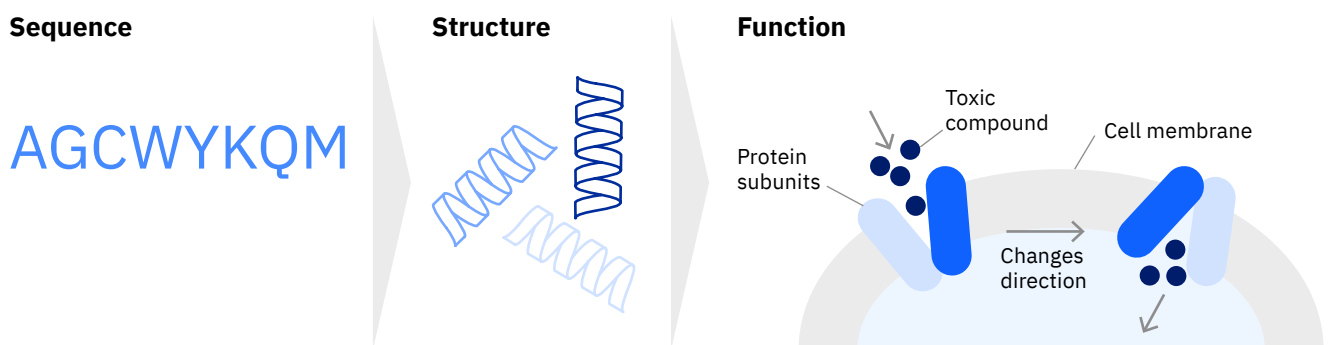
exhaustively searching all possible arrangements for a given length. New insights from quantum computing could further our understanding of transcription factor binding and de novo genome assembly.

- *Genome-wide association studies (GWAS)*<sup>10</sup>: The goal of GWAS is to find associations between a selected trait or disease and single mutations in DNA. Current methods are inherently high-dimensional and computationally challenging. This highlights the potential for quantum computing, which may significantly narrow the typically long lists of candidate genes that need to be experimentally validated. Quantum computing may also enable progress in gene network and graph models.
- *De novo structure prediction*<sup>11</sup>: With the explosive growth of sequencing information and technology, an increasing gap exists in understanding how sequence translates to structure and, ultimately, function (see Figure 2). Despite sophisticated methods, such as homology models, classical approaches to predict structure de novo often scale poorly.<sup>12</sup> For instance, the search space of potential protein configurations increases exponentially with the size of the protein (see Use Case 3). Quantum computing has the potential to drastically improve structure predictions for RNA molecules, proteins, DNA-protein complexes, and other constructs.

—

**Figure 2**

Sequence-structure-function dogma, which is at the heart of biology research



## Quantum computing has a diversity of potential applications in drug discovery.

Such advances could eventually help realize the vision of powerful digital twin models.<sup>13</sup> Organic digital twins might be used in pharmacogenomic testing to predict an individual's response to specific drugs over time, aiding the development of precision medicine therapies. Additional inorganic digital twins could be created to optimize research or care facilities by comparatively stress-testing aspects such as procedures, staffing levels, facility layout, and equipment. Reaching the day when a medical team can tell a patient, "Based on your genome, we have confidence that this will be the specific result of your treatment," may someday no longer seem like a purely utopian goal.

### **Use Case 2: Improving patient outcomes by enhancing the efficiency of small-molecule drug discovery**

Small-molecule drug design and discovery has always been a complex optimization process. Its goal: improving patient outcomes by designing a novel molecule active against the disease-related target while simultaneously reducing activity against the thousands of other targets in the body to avoid side effects and dangerous toxicities. In pursuit of this goal, typically 200,000 to  $>10^6$  compounds are screened in experimental and computational workflows, and a few thousand are produced and tested in the necessary battery of assays.<sup>14</sup> Here, computing has long played a role, largely through similarity and classification approaches to support screening and detailed 3D structure, as well as energy calculations to support more precise target-based design.

Quantum computing has a diversity of potential applications in drug discovery.<sup>15</sup> The technology could help assess a greater number of candidate molecules and evaluate them more accurately using, for example, classification methods such as those employed in

lead-finding and off-target screening. And it may impact the classification associated with the high-throughput task of lead-finding and the modeling of off-targets in lead optimization—as well as with the physics-based modeling carried out in lead optimization when a 3D protein structure or good model is available.

The ability to explore more molecules is important since the number of possible small molecules is enormous. Normally only a small fraction is considered. In fact, the total number of possible carbon-based compounds whose molecular masses are similar to those of living systems is around  $10^{60}$ .<sup>16</sup> Effectively exploring this chemical space is an area of great potential. It opens the door to better assessing ultra-large libraries of small organic molecules now available for purchase with synthesis "on demand."<sup>17</sup>

Particularly accurate scoring is possible through molecular dynamics simulations of protein-ligand complexes. Here, quantum computing could offer significant advantages for carrying out hybrid quantum/molecular mechanics applications as well as developing the underlying parameters of the classical force field. Such advances would apply to both lead optimization as well as the growing field of computational process chemistry, such as in modeling enzymatic reactivity and stereoselectivity to support biocatalysis in drug manufacturing.<sup>18</sup>

### **Use Case 3: Developing novel biological products based on protein folding predictions**

In contrast to small-molecule drugs, in the case of biologics, a protein or other macromolecule is the drug. Biological drugs, such as antibodies, insulin, and many vaccines, have been employed for decades.<sup>19</sup> In recent years, pharmaceutical companies are increasingly targeting biologics to treat a number of diseases. Designing the 3D structure of biologics is important for function, specificity, and stability.<sup>20</sup>

Real-world protein modeling cases involve exploration of the enormous number of possible folding patterns, as illustrated in Levinthal’s paradox (see Figure 3).<sup>21</sup> The exponential growth of potential conformations with chain length makes the problem challenging for classical computers. For example, in one model, a chain of 20 amino acids has  $10^9$  potential conformations, and chains with 60 and 100 amino acids have  $10^{28}$  and  $10^{47}$  conformations, respectively.<sup>22</sup> Moreover, as part of the FDA’s biological product definition, a protein must comprise more than 40 amino acids.<sup>23</sup>

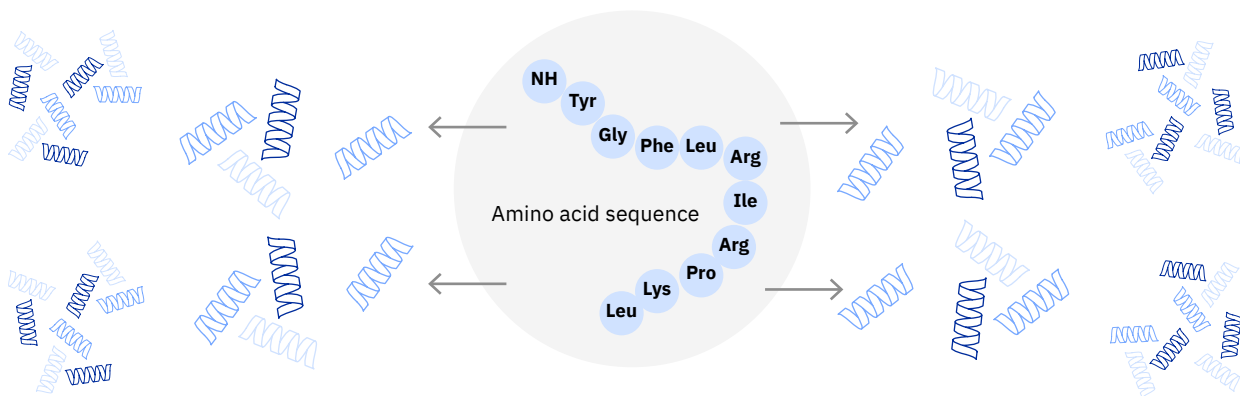
While many proteins can be modeled adequately by analogy to known structures, an important and challenging design target is the hypervariable H3 loop in the complementarity-determining region of antibodies.

This loop typically contains 3–20 residues, but is sometimes much longer, and accurate representation has been the subject of much study.<sup>24</sup>

Quantum computing has the potential to overcome many of these computational challenges—for example, scoring the great number of possible structures and identifying the likeliest one. A recent publication demonstrated that quantum computing could score a peptide in two common conformations represented on a lattice—alpha helix and beta sheet—and leveraged a quantum algorithm for the search.<sup>25</sup> It has also been shown that quantum computing may drastically improve the calculation of protein force fields.<sup>26</sup> As quantum volume increases, quantum computing’s ability to score additional conformations will increase accordingly.<sup>27</sup>

**Figure 3**

Levinthal’s paradox – even a protein with only 100 amino acids has around  $10^{47}$  potential conformations. In reality, however, many proteins fold to their native structure within seconds.



Finally, as with all the potential quantum applications discussed in this report, quantum computing could enable further use cases in tangential areas. For instance, biologics tend to be much less stable than small-molecule drugs. Optimization of the biologics supply chain itself—from formulation, through shipment, and ultimately transport to pharmacies, hospitals, and even homes—is a complicated process that also may be improved by quantum computing.<sup>28</sup>

## From bench to bedside

The life sciences sector has the potential to benefit significantly from quantum computing. Trends, such as the spread of efficient low-cost sequencing and the advent of the “-omics” era, result in life sciences companies exploring ways to take advantage of this diversity of novel data sources. Further, it is among the industries in which people could most directly experience future quantum computing benefits.

Exploration and implementation of quantum use cases, paired with further scientific progress in quantum hardware and algorithms, is expected to enable the transition from potential to reality over the coming years. Leading life sciences and pharmaceutical companies have started this journey toward quantum advantage.

## Action guide

### *Exploring quantum computing use cases for life sciences*

Quantum computing necessitates a different way of thinking, a new and highly sought-after set of skills, distinct IT architectures, and novel corporate strategies. Imagine some of the capabilities and benefits described in this report in the hands of your top competitors, particularly as much of the early IP in quantum computing may be proprietary. That’s why the time to get started with quantum computing is now—when standards, strategies, use cases, and ecosystems are being developed.

So how can life sciences organizations get started with quantum computing? There are three key initial steps:

**1. Identify and enable quantum champions in your organization to experiment with actual quantum computers and explore potential applications for your specific industry.** To help focus on your highest-value problems, have your quantum champions report to a quantum steering committee that includes line-of-business executives, innovation and technology leads, and market strategists.

**2. Prioritize quantum use cases according to their potential for attaining business advantage—given your organization’s therapeutic focus areas, business strategy, associated customer value propositions, and future growth plans.** Keep an eye on progress in quantum application development to stay in the vanguard of which use cases might be commercialized sooner rather than later.

**3. Consider partnering with an emerging quantum ecosystem of like-minded research labs and academic institutions, technology providers, application developers, and start-ups.** Obtain access to an entire quantum computing stack capable of developing and running quantum algorithms specific to your business needs. Look for breakthroughs in quantum technology that might necessitate a change in ecosystem partners.



## Notes and sources

- 1 Bohacek, Regine S., Colin McMartin, Wayne C. Guida. "The art and practice of structure-based drug design: A molecular modeling perspective." *Medicinal Research Reviews*. January 1996. [https://azevedolab.net/resources/The%20art%20and%20practice%20of%20structure-based%20drug%20design\\_%20A%20molecular%20modeling%20perspective.pdf](https://azevedolab.net/resources/The%20art%20and%20practice%20of%20structure-based%20drug%20design_%20A%20molecular%20modeling%20perspective.pdf)
- 2 Feynman, Richard P. "Simulating Physics with Computers." *International Journal of Theoretical Physics*. May 7, 1981. <https://catonmat.net/ftp/simulating-physics-with-computers-richard-feynman.pdf>
- 3 "5in5: Five innovations that will change our lives within five years." IBM. 2019. <https://www.research.ibm.com/5-in-5/quantum-computing>
- 4 Havlíček, Vojtěch, Antonio D. Córcoles, Kristan Temme, Aram W. Harrow, Abhinav Kandala, Jerry M. Chow, Jay M. Gambetta. "Supervised learning with quantum-enhanced feature spaces." *Nature*. March 13, 2019. <https://arxiv.org/pdf/1804.11326.pdf>; Carleo, Giuseppe, Matthias Troyer. "Solving the Quantum Many-Body Problem with Artificial Neural Networks." *Science*. February 10, 2017. <https://arxiv.org/pdf/1606.02318.pdf>
- 5 Montanaro, Ashley. "Quantum algorithms: an overview." *Nature*. January 12, 2016. <https://www.nature.com/articles/npjqi201523>
- 6 Tirrell, Meg. "Unlocking my genome: Was it worth it?" *CNBC*. December 14, 2015. <https://www.cnn.com/2015/12/10/unlocking-my-genome-was-it-worth-it.html>; Kandpal, Raj P., Beatrice Saviola, Jeffrey Felton. "The era of 'omics unlimited.'" *Future Science*. April 25, 2018. <https://www.future-science.com/doi/full/10.2144/000113137>
- 7 Copeland, Libby. "You Can Learn a Lot About Yourself From a DNA Test. Here's What Your Genes Cannot Tell You." *Time*. March 2, 2020. <https://time.com/5783784/dna-testing-genetics>
- 8 Emani, Prashant S., Jonathan Warrell, Alan Anticevic, Stefan Bekiranov, Michael Gandal, Michael J. McConnell, Guillermo Sapiro, Alán Aspuru-Guzik, Justin Baker, Matteo Bastiani, Patrick McClure, John Murray, Stamatios N. Sotiropoulos, Jacob Taylor, Geetha Senthil, Thomas Lehner, Mark B. Gerstein, Aram W. Harrow. "Quantum Computing at the Frontiers of Biological Sciences." 2019. <https://arxiv.org/ftp/arxiv/papers/1911/1911.07127.pdf>
- 9 Zambelli, Federico, Graziano Pesole, Giulio Pavesi. "Motif discovery and transcription factor binding sites before and after the next-generation sequencing era." *Briefings in Bioinformatics*. April 19, 2012. <https://academic.oup.com/bib/article/14/2/225/208333>
- 10 "Genome-Wide Association Studies Fact Sheet." National Human Genome Research Institute. August 27, 2015. <https://www.genome.gov/about-genomics/fact-sheets/Genome-Wide-Association-Studies-Fact-Sheet>
- 11 Das, Rhiju, David Baker. "Automated de novo prediction of native-like RNA tertiary structures." *PNAS*. September 11, 2007. <https://www.pnas.org/content/pnas/104/37/14664.full.pdf>
- 12 Muhammed, Muhammed Tilahun, Esin Aki-Yalcin. "Homology modeling in drug discovery: Overview, current applications, and future perspectives." *Chemical Biology & Drug Design*. September 6, 2018. <https://onlinelibrary.wiley.com/doi/full/10.1111/cbdd.13388>

- 13 Fuller, Aidan, Zhong Fan, Charles Day, Chris Barlow. "Digital Twin: Enabling Technology, Challenges and Open Research." *Deep AI*. October 29, 2019. <https://arxiv.org/pdf/1911.01276.pdf>
- 14 Hughes, JP, S Rees, SB Kalindjian, KL Philpott. "Principles of early drug discovery." *British Journal of Pharmacology*. November 22, 2010. <https://bpspubs.onlinelibrary.wiley.com/doi/full/10.1111/j.1476-5381.2010.01127.x>
- 15 Cao, Yudong, Jhonathan Romero, Alán Aspuru-Guzik. "Potential of quantum computing for drug discovery." *IBM Journal of Research and Development*. November –December 1, 2018. <https://ieeexplore.ieee.org/document/8585034>
- 16 Dobson, Christopher M., "Chemical space and biology." *Nature*. December 15, 2004. <https://www.nature.com/articles/nature03192>
- 17 Lyu, Jiankun, Sheng Wang, Trent E. Balius, Isha Singh, Anat Levit, Yurii S. Moroz, Matthew J. O'Meara, Tao Che, Enkhjargal Algaa, Kateryna Tolmacheva, Andrey A. Tolmachev, Brian K. Shoichet, Bryan L. Roth, John J. Irwin. "Ultra-large library docking for discovering new chemotypes." *Nature*. February 6, 2019. <https://www.nature.com/articles/s41586-019-0917-9>
- 18 Cao, Yudong, Jonathan Romero, Jonathan P. Olson, Matthias Degroote, Peter D. Johnson, Maria Kieferova, Ian D. Kivlichan, Tim Menke, Borja Peropadre, Nicolas P. D. Sawaya, Sukin Sim, Libor Veis, Alan Aspuru-Guzik. "Quantum Chemistry in the Age of Quantum Computing." *Chemical Reviews*. August 30, 2019. <https://arxiv.org/pdf/1812.09976.pdf>
- 19 Middaugh, C.R., R. Pearlman. "Proteins as Drugs: Analysis, Formulation and Delivery." *Novel Therapeutics from Modern Biotechnology. Handbook of Experimental Pharmacology*, vol 137. [https://link.springer.com/chapter/10.1007/978-3-642-59990-3\\_3](https://link.springer.com/chapter/10.1007/978-3-642-59990-3_3)
- 20 Johnston, Sarah L. "Biologic therapies: what and when?" *Journal of Clinical Pathology*. March 2007. <https://www.ncbi.nlm.nih.gov/pmc/articles/PMC1860592>
- 21 Levinthal, Cyrus, "How to fold gracefully." *Mössbaun Spectroscopy in Biological Systems Proceedings*. 1969. [https://www.cc.gatech.edu/~turk/bio\\_sim/articles/proteins\\_levinthal\\_1969.pdf](https://www.cc.gatech.edu/~turk/bio_sim/articles/proteins_levinthal_1969.pdf)
- 22 Zanzig, Robert, Attila Szabo, Biman Bagchi. "Levinthal's paradox." *Proceedings of the National Academy of Science*. October 7, 1991. <https://www.pnas.org/content/pnas/89/1/20.full.pdf>
- 23 Mezher, Michael. "FDA Finalizes 'Biological Product' Definition Ahead of BPCIA Transition." *Regulatory Focus*. February 20, 2020. <https://www.raps.org/news-and-articles/news-articles/2020/2/fda-finalizes-biological-product-definition-aea>
- 24 Marks, C., C.M. Deane. "Antibody H3 Structure Prediction." *Computational and Structural Biotechnology Journal*. January 24, 2017. <https://www.sciencedirect.com/science/article/pii/S2001037016301118>
- 25 Robert, Anton, Panagiotis Kl. Barkoutsos, Stefan Woerner, Ivano Tavernelli. "Resource-Efficient Quantum Algorithm for Protein Folding." August 6, 2019. <https://arxiv.org/pdf/1908.02163.pdf>
- 26 Mishra, Anurag, Alireza Shabani. "High-Quality Protein Force Fields with Noisy Quantum Processors." October 29, 2019. <https://arxiv.org/pdf/1907.07128.pdf>
- 27 Chow, Jerry, Jay Gambetta. "Quantum Takes Flight: Moving from Laboratory Demonstrations to Building Systems." *IBM*. January 8, 2020. <https://www.ibm.com/blogs/research/2020/01/quantum-volume-32>
- 28 "Transparency19: The future of freight is here." *Shipwell*. May 14, 2019. <https://shipwell.com/company-news/transparency-2019-wrap-up>

## About Expert Insights

Expert Insights represent the opinions of thought leaders on newsworthy business and related technology topics. They are based upon conversations with leading subject matter experts from around the globe. For more information, contact the IBM Institute for Business Value at [iibv@us.ibm.com](mailto:iibv@us.ibm.com).

© Copyright IBM Corporation 2020

IBM Corporation  
New Orchard Road  
Armonk, NY 10504

Produced in the United States of America  
April 2020

IBM, the IBM logo, and [ibm.com](http://ibm.com) are trademarks of International Business Machines Corp., registered in many jurisdictions worldwide. Other product and service names might be trademarks of IBM or other companies. A current list of IBM trademarks is available on the web at “Copyright and trademark information” at [www.ibm.com/legal/copytrade.shtml](http://www.ibm.com/legal/copytrade.shtml).

This document is current as of the initial date of publication and may be changed by IBM at any time. Not all offerings are available in every country in which IBM operates.

THE INFORMATION IN THIS DOCUMENT IS PROVIDED “AS IS” WITHOUT ANY WARRANTY, EXPRESS OR IMPLIED, INCLUDING WITHOUT ANY WARRANTIES OF MERCHANTABILITY, FITNESS FOR A PARTICULAR PURPOSE AND ANY WARRANTY OR CONDITION OF NON-INFRINGEMENT. IBM products are warranted according to the terms and conditions of the agreements under which they are provided.

This report is intended for general guidance only. It is not intended to be a substitute for detailed research or the exercise of professional judgment. IBM shall not be responsible for any loss whatsoever sustained by any organization or person who relies on this publication.

The data used in this report may be derived from third-party sources and IBM does not independently verify, validate or audit such data. The results from the use of such data are provided on an “as is” basis and IBM makes no representations or warranties, express or implied.

