

DATA-DRIVEN SEARCH AND INNOVATION *

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Preliminary Draft – Please Do Not Circulate

First Version: June 12th, 2021

This Version: February 5th, 2023

Abstract

In a growing variety of contexts, correlational data are being used to find successful innovations even when the inventors do not fully grasp why they should work. This method of data-driven search for innovation stands in stark contrast with traditional approaches that rely on abstract knowledge to discover breakthroughs. In this project, I develop a novel theoretical framework to explain how data changes the recombinant search for innovation and discuss implications in terms of search breadth and breakthrough discovery. Next, I empirically investigate this phenomenon in the context of human genomics, where the advent of genome-wide association studies (GWASs) approximates the ideal of a data-driven search for the genetic roots of diseases. By comparing gene-disease associations introduced by GWASs with those from theory-driven studies, I provide unique evidence of how the search process shapes innovation outcomes. My results show that gene-disease associations introduced by GWASs span a wider portion of the genetic landscape, are more likely to involve neglected human genes, and are of higher scientific value than comparable associations introduced by theory-driven studies. However, heterogeneity analyses reveal that data-driven search performs poorly with interdependent genes because correlational data neglect complex interactions that only a theoretical understanding can capture. This paper contributes to our understanding of how data change innovation and the conditions under which they might unlock breakthrough discoveries.

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1 Introduction

Innovation is generated by recombining technological components in new ways. The universe of potential combinations is often depicted as a landscape over which innovators search (Aharonson and Schilling, 2016; Fleming and Sorenson, 2001). Such landscape has peaks of value in correspondence with the best opportunities but also large valleys of low-value recombinations. Previous research has documented a proclivity of individuals and organizations to over-focus on incremental modifications of past successful combinations (Audia and Goncalo, 2007; Rzhetsky et al., 2015; Stuart and Podolny, 1996). But when search in familiar domains necessarily runs into technological exhaustion, exploration of uncharted spaces becomes necessary to discover new peaks (March, 1991; Levinthal, 1997; Fleming, 2001). This realization leads to one of the crucial questions in management research: how can innovators find breakthrough recombinations in unknown technological spaces?

A growing body of research has suggested that innovators should be guided by abstract reasoning and cause-effect knowledge (Arora and Gambardella, 1994; Ehrig and Schmidt, 2022; Fleming and Sorenson, 2004). By developing theoretically-motivated hypotheses on which combinations should work, inventors can find the best innovations more efficiently compared to haphazard trial-and-error (Camuffo et al., 2022; Gavetti and Levinthal, 2000; Felin and Zenger, 2017; Kneeland et al., 2020). However, this answer seems at odds with how many successful firms search nowadays (Allen, 2022; Agrawal et al., 2022; Thomke, 2020). For instance, scientists in pharmaceutical firms gather information on millions of compounds, without knowing *ex ante* which ones might work as drugs (Evans and Rzhetsky, 2010; Jayaraj and Gittelman, 2018). Entrepreneurs use A/B testing to triage ideas at scale, iteratively testing many configurations without assuming why some should be better (Koning et al., 2022). In an increasing variety of fields, innovators leverage large quantities of data instead of their theoretical priors to find peaks in novel domains (Nagaraj, 2022).

In this paper, I theorize that the ability to generate and analyze “big data” offers a new way to innovate in unknown technological spaces. Large-scale data allows the extraction of signals on the value of technological combinations, an approach that I call *data-driven search*. Inventors can use data to search through vast combinatorial spaces and triage potential combinations at scale, without relying on theoretical priors or costly experiments (Agrawal et al., 2022; Choudhury et al., 2021; Christin, 2020; Shrestha et al., 2020). While theories operate by reducing the dimensionality of search problems (Gavetti and Levinthal, 2000), thus narrowing down search to areas that seem promising (Fleming and Sorenson, 2004), data-driven search enlarges the scope of combinations that can be assessed offline. Relative to theory-driven search, the result should be an increase

in search breadth that might reduce the tendency to focus only on well-understood areas of the combinatorial landscape.

While data enables a novel strategy for recombinant search, its effect on the quality of innovation is *ex ante* ambiguous (Schilling and Green, 2011). On the one hand, data-driven search should facilitate recombinations from distant technological domains because it is not bound to components theoretically understood (Anderson, 2008; Wu et al., 2020). Broadening the search scope could spur creativity and outperform theory-driven search if its focus is too narrow (Katila and Ahuja, 2002; Leiponen and Helfat, 2010). On the other hand, abstract knowledge of cause-effect mechanisms allows recombining technological elements more effectively (Arora and Gambardella, 1994; Arts and Fleming, 2018; Nelson, 1982). Moreover, if agents are already searching in the most fruitful areas of the landscape, then increasing the breadth of search will not improve the value of innovations (Fleming and Sorenson, 2004; Kaplan and Vakili, 2015; Rzhetsky et al., 2015). Overall, whether and under what conditions data-driven search will prove more effective at locating breakthrough combinations than theory-driven search is an empirical question.

Empirical investigation of these ideas requires comparing theory-driven and data-driven search strategies. This is challenging for two reasons. First, one must find a well-defined search problem involving observable technological components. In particular, a meaningful comparison necessitates observing both search strategies used in the same landscape, on the same recombinatorial task, and by comparable actors. Second, one needs to characterize distinct search processes and be able to tie them with the resulting innovation outputs. This is especially difficult because the researcher can only see realized outcomes without usually being able to know what kind of search process generated them (Schilling and Green, 2011; Kneeland et al., 2020; Maggitti et al., 2013). Investigating the consequences of data-driven search requires finding a setting where all these conditions are met at the same time.

In this paper, I address both challenges with an empirical study of how scientists search for the genetic roots of human diseases. First, genes and diseases constitute the relevant components for this search problem. Any gene could be tied to any condition, generating a space of millions of potential gene-disease combinations. The key objective of researchers is finding which combinations can serve as targets for drug development. Second, searching for gene-disease associations can happen in two ways. Scientists interested in a disease can carry out *candidate gene studies* using their theoretical priors to target specific genes, or they can perform atheoretical *genome-wide association studies* (GWASs) that scan the whole genome to locate gene variants correlated to the disease. Importantly, I can infer what search process led to a specific gene-disease combination by coding

the method used in the scientific article that introduced it. Comparing combinations established by GWASs to those discovered by candidate gene studies allows me to descriptively explore the characteristics and impact of data-driven search in this task so crucial for drug development.

I assemble a new dataset that includes all gene-disease associations (GDAs) introduced in the period 1980-2016. The raw data are taken from DisGeNET, the most comprehensive aggregator of information on the genes associated with human diseases. DisGeNET collects the list of all PubMed articles that studied each GDA, including the publication that first originated it and the list of all subsequent papers that investigated it. The count of follow-on studies that directly explored a gene-disease association provides a precise measure of its scientific impact, regardless of whether such studies cite the paper that first introduced it. Next, I use data from the European Bioinformatics Institute to identify papers that use a genome-wide approach, thus being able to code which associations were established by a GWAS and which ones by candidate gene approaches. I empirically compare theory-driven and data-driven search on two related dimensions: the direction of search efforts and the scientific potential of the gene-disease combinations uncovered.

My empirical analysis suggests that data-driven search diversifies innovation by exploring a wider portion of the genetic landscape. In baseline estimates, gene-disease combinations discovered by GWASs are 114-155% and 86-108% more likely to involve genes understudied or recently discovered, respectively. Additional tests suggest this is because genome-wide association studies remove the ex ante choice of what genes to target, hence overcoming the path dependency that characterizes theory-driven search. Further, a comparison between data-driven and theory-driven search shows that the former discovers more breakthrough gene-disease combinations even holding the characteristics of genes and diseases constant. To mitigate endogeneity concerns, I include principal investigator fixed effects, thus effectively controlling for time-invariant characteristics of the researcher that might correlate with the ability to find breakthrough discoveries. Even among gene-disease associations introduced by the same researcher, those uncovered with a genome-wide association study are 77.4% more likely to be in the top tail of scientific impact.

Heterogeneity analyses show that the beneficial effects of data-driven search are contingent on the area of the landscape explored. I find that data are beneficial to locate the best opportunities in uncharted technological areas, where there is little theoretical guidance on how genes and diseases are related. However, candidate gene studies largely outperform GWAS in areas with a better theoretical understanding of genetic biology. To address the concern that the distribution of theoretical knowledge on the landscape is endogenous (Gittelman, 2016; Nelson, 2003), I exploit the fact that certain genes have historically been neglected simply because they cannot be studied

using the lab mouse (Stoeger et al., 2018). Exploiting this variation in theory availability unrelated to the therapeutic potential of the genes, I confirm that data-driven search can effectively locate breakthroughs among technological components that are less theoretically known.

Further investigation of the mechanisms behind these results shows that genome-wide findings are less impactful when they involve genes that regulate the function of multiple downstream genes (Hermosilla and Lemus, 2019). This suggests a fundamental limitation of data-driven search: data signals report correlations that are unable to consider complex interactions between interdependent or coupled components (Fleming and Sorenson, 2004; Ghosh, 2021; Kapoor and Wilde, 2022). Instead, in the presence of a solid-enough theoretical understanding, researchers can locate the best gene-disease combinations because they can take into account how complex genes interact. Additional robustness tests further confirm that my results are not due to the characteristics of scientists adopting data-driven search, the choice of diseases studied, or the specific definition of breakthrough employed.

This work contributes to understanding the mechanisms through which data are reshaping innovation. I theorize that data enable an alternative way to search for valuable recombinations that does not rely on theories or cause-effect understandings. I provide empirical evidence that data allow to expand the search scope, increasing the ability to recombine technological components that are overlooked because they are less familiar to the inventor. While these findings are descriptive and limited to a specific (albeit important) search problem, they suggest that data can increase breakthrough innovation by expanding search breadth. The heterogeneity results indicate that managers and inventors can rely on data when exploring uncharted domains but should be wary that correlational leads fail to take into account complex interdependencies. When dealing with interdependent components, data signals can be a misleading guide, raising the value of theoretical knowledge.

The paper proceeds as follows. Section 2 describes the construct of data-driven search and discusses its implications for discovering breakthrough innovation. Section 3 provides an overview of how scientists search for gene-disease associations and explains the characteristics of genome-wide association studies. Section 4 describes the data and research design, while Section 5 presents empirical estimates of the role of data in shaping search patterns and breakthrough innovation. Section 6 concludes.

2 Theoretical Framework

2.1 Data-Driven Search

Recent years have seen the emergence and diffusion of large-scale datasets. Researchers have started investigating their impact on corporate decision-making (Brynjolfsson and McElheran, 2016), organizational structure (Wu et al., 2019), and scientific production (Nagaraj et al., 2020; Nagaraj and Tranchero, 2022). However, when focusing on innovation, the results are ambiguous and heterogeneous (Deniz, 2020; Brynjolfsson et al., 2021; Ghosh, 2021; Lou and Wu, 2021; Wu et al., 2020). These mixed findings appear at odds with the hype surrounding big data and underscore the need for a better understanding of how data are actually used in technological innovation.

In this paper, I propose that the availability of large-scale data unlocks a new strategy to search for promising technological combinations. Instead of relying on knowledge from past attempts or theoretical priors, inventors can use data to extract signals of what combinations seem more fruitful for follow-on experimentation, a process that can be described as *data-driven search*. In practice, data can be used to triage and rank potential combinations *in silico*, thus focusing on the most promising ones without the need for actual experimentation (Christin, 2020; Hoelzemann et al., 2022). The power of this approach is that data can help assessing a vast number of combinations that would be impossible to individually try out or theoretically model (Choudhury et al., 2021; Evans and Rzhetsky, 2010; Shrestha et al., 2020). Moreover, innovators no longer need to pre-select a subset of components to recombine based on their prior knowledge, a choice that leads to path dependency in search (Gavetti and Levinthal, 2000; Evans, 2010; Rzhetsky et al., 2015; Stuart and Podolny, 1996).

Data-driven search requires a few conditions to be feasible. First, the relevant characteristics of the technological components must be measurable, ensuring that the space of possible combinations is well-defined. This means that data-driven search might be of little help when trying to invent entirely new technologies that do not emerge from old components (Wu et al., 2020). Second, there has to exist a metric of technological potential on which the promise of each combination can be assessed. Such metric constitutes the objective function that data-driven search tries to maximize by finding the candidate combinations that score highest (Agrawal et al., 2022). Third, and relatedly, it must be possible to foresee the effect of novel combinations on the objective of interest. Said otherwise, data-driven search requires that it is possible to predict the value of potential recombinations from the data available on components. Appendix A presents an example

from combinatorial chemistry that illustrates how these boundary conditions define the feasibility of data-driven search.

Given the conditions outlined above, it seems that data-driven search happens mainly within the boundaries of the prevailing technological and scientific paradigms (Dosi, 1982; Kuhn, 1962). In turn, this has two main implications. First, the paradigm can help determining the metric used to assess the value of alternative technological combinations from data signals. This means that meta-theoretical understandings of “what is important” will concur in shaping the ability to locate breakthrough innovations. Second, the decision of where to direct the data-collection effort might itself reflect the priorities implicit in the current paradigm. Insofar as this choice is biased, it might end up reinforcing previously established research patterns (Cao et al., 2021; Hoelzemann et al., 2022). However, the increasing diffusion of complete maps of diverse technological landscapes, such as the Sloan Digital Sky Survey or the Human Genome Project, should reduce the practical relevance of this concern (Nagaraj and Stern, 2020).

2.2 How Does Data Change Recombinant Search?

Comparing data-driven search with alternative approaches to uncover breakthroughs can help to discern how data reshape innovation. In particular, a strand of work has suggested that innovators should leverage their theoretical priors to focus search on the portions of the combinatorial space expected to yield the highest returns (Csaszar and Levinthal, 2016; Gavetti and Levinthal, 2000; Klahr, 2000; Nelson, 1982). In turn, the source of inventors’ priors can range from scientific information (Fleming and Sorenson, 2004; Gambardella, 1995; Kneeland et al., 2020) to abstract theories of cause-effect relationships among components (Arora and Gambardella, 1994; Ehrig and Schmidt, 2022; Felin and Zenger, 2017). Inventors experiment with combinations that have the highest expected value based on their current beliefs (Camuffo et al., 2022), following a *theory-driven search* (Gittelman, 2016).

Gavetti and Levinthal (2000) note that search strategies can be characterized by three properties: how alternatives are evaluated, the extensiveness of other options considered, and how distant they are from the area currently being searched. Starting with the mode of evaluation, both theory-driven and data-driven search operate *off-line*, meaning that they assess the merit of alternative combinations without the need to actually invest in experimentally validating them. However, they diverge in the scope of alternatives considered (Agrawal et al., 2022). Any theory, however accurate, can only account for some dimensions of the reality represented (Gavetti and Levinthal, 2000). Reducing the dimensionality of search problems will necessarily come at the cost of narrowing

search on some dimensions. In contrast, big data allow obtaining signals about a potentially much larger set of alternative combinations.

In turn, the crucial difference in the extensiveness of alternatives considered will likely shape the location of innovative activities. Theory-driven search exploits known cause-effect relationships to funnel experimentation efforts in areas where inventors theoretically understand how components could be recombined (Arora and Gambardella, 1994). The result is usually a persistent search in the same narrow portion of the landscape, even in the face of decreasing returns (Gavetti and Levinthal, 2000; Fleming and Sorenson, 2004; Helfat, 1994). Instead, data-driven search breaks the path dependency that characterizes theory-driven search (Helfat, 1994; Rzhetsky et al., 2015). Extracting signals from data should not be biased by beliefs or capabilities that steer innovators away from explorative attempts (March, 1991), unless the data themselves are fundamentally biased (Cao et al., 2021). As a result of greater search breadth, innovators will be more likely to recombine components further away from domains previously explored (Katila and Ahuja, 2002; Wu et al., 2020).

2.3 The Consequences of Data-Driven and Theory-Driven Search

Does data-driven search lead to discovering more breakthrough innovations relative to theory-driven search? Existing research does not offer a clear-cut answer. Theories tend to focus efforts where returns are expected to be highest, possibly at the cost of neglecting areas that might harbor breakthroughs (Gavetti and Levinthal, 2000; Klahr, 2000; Langley et al., 1987). Instead, novel discoveries might require going against established beliefs to bridge diverse knowledge (Audia and Goncalo, 2007; Fleming, 2001). Data-driven search could be useful in such uncharted domains because data signals help triage recombinations even if the mechanisms of what makes a combination valuable are unknown (Gambardella, 1995; Gittelman, 2016; Wu et al., 2020). Insofar as breakthrough innovations are the result of bridging distant technological domains, we should then expect data-driven search to yield more breakthroughs than theory-driven approaches (Schilling and Green, 2011).

However, it could also be that reliance on data affects search in unwarranted ways. The literature has noted that innovators face a trade-off between depth and breadth of search, because cumulative learning can be essential to generate novelty (Helfat, 1994; Leiponen and Helfat, 2010; Katila and Ahuja, 2002; Stuart and Podolny, 1996). Atheoretical exploration of diverse domains might impede value identification because one needs sufficient knowledge to assess data signals about unconventional ideas (Kaplan and Vakili, 2015; Lou and Wu, 2021; Wu et al., 2020). On the

contrary, a deep understanding of an area can provide the necessary capabilities to find and unlock breakthrough recombinations (Arts and Fleming, 2018; Fleming and Sorenson, 2004; Nelson, 1982). Moreover, if innovators are already searching in the most fruitful areas of the landscape, then increasing the breadth of search will not increase breakthroughs (Fleming and Sorenson, 2004).

Considering these arguments, whether data-driven search will prove more effective at locating breakthrough combinations than theory-driven search is an empirical question. However, research on recombinant search suggests that the complexity of the combinatorial task is a key moderator of search effectiveness (Aharonson and Schilling, 2016; Fleming and Sorenson, 2001). Theories should be especially useful to foresee the value of combinations between highly interdependent elements (Fleming and Sorenson, 2004). Data signals might offer poor guidance in those instances since they can, at best, provide correlations that are unable to account for complex interactions between components (Deniz, 2020; Ghosh, 2021). Therefore, in light of the above discussion, one can hypothesize that the effectiveness of either search strategy will be contingent on the area of the landscape explored. In the remainder of the paper, I empirically assess this idea by examining how data are used to search for the genetic roots of human diseases.

3 Empirical Setting

3.1 Scientific Background

Genes are sequences of DNA bases that encode the “instructions” to synthesize gene products with a fundamental role in the organism’s functioning. Knowing the genetic roots of diseases has significant practical consequences since genes causative of diseases can potentially serve as drug targets (Nelson et al., 2015). Most common diseases such as diabetes, Alzheimer’s, or hypertension are *polygenic*: they are not due to a single genetic factor but rather to multiple genes and their interaction with the environment during human life (Bush and Moore, 2012). Discovering the genes involved in each of the thousands of polygenic diseases requires searching through the $\sim 20,000$ known human genes. How do scientists look for new gene-disease associations in this huge combinatorial space?

Scientists traditionally followed a *candidate gene approach* consisting of three main steps (Tabor et al., 2002). First, the scientist decides the disease to study, likely motivated by its prevalence or funding availability. Second, she hypothesizes what genes might have a role in its etiology. Finally, she focuses the analysis on those genes, typically employing family linkage studies, case-control studies, or experiments with lab animals. Importantly, selecting the target genes reflects scientists’

biological understanding of which genes might be important and why. For instance, after the IL12B gene was associated with psoriasis, some scientists relied on their knowledge of IL12B's role in the metabolic pathway of the IL23R gene to hypothesize a connection between IL23R and psoriasis. Indeed, this reasoning led to the candidate-gene study that first documented the role of IL23R in psoriasis (Cargill et al., 2007).

Despite several successful discoveries, candidate gene studies have led scientists to consider only genes well known theoretically (Haynes et al., 2018). The result has been an extreme concentration of attention on a small number of genes (Oprea et al., 2018; Stoeger et al., 2018). Gates et al. (2021) report that 1% of the genome has received 22% of all gene-related publications. This situation is suboptimal since our understanding of polygenic diseases would benefit from exploring a larger pool of genes (Edwards et al., 2011). The excessive research emphasis on a handful of heavily studied genes means that researchers ignore a large number of potentially important genes (Stoeger et al., 2018). As a consequence, currently approved drugs exploit only around 10% of the potential drug targets highlighted by the Human Genome Project, leaving many therapeutic opportunities still untried (Gates et al., 2021)

3.2 Genome-Wide Association Studies as Data-Driven Search

Starting from the early 2000s, two events concurred in providing an alternative to candidate gene studies. The first was the completion of the International HapMap Project in 2005. The HapMap was designed to provide a detailed reference genome that could be used to relate genetic mutations with phenotype changes (Bush and Moore, 2012). The second, and related, was the diffusion of commercial genotyping microarrays. Unlike whole genome sequencing, which processes every DNA basis, microarrays only detect the activity of specific genetic loci. The HapMap enabled to design microarrays targeting loci that can be extrapolated to capture the characteristics of their genetic surroundings, thus allowing to parsimoniously infer the characteristics of most of the genome (Bush and Moore, 2012). The result was a steep decrease in the cost of collecting data on genomes, prompting the emergence of *genome-wide association studies* (Visscher et al., 2017).

Genome-wide association studies (or GWASs) are case-control studies where researchers sequence a large number of genomes and look to see if any genetic variation is more likely to appear in subjects showing a specific condition rather than in the control group (Pearson and Manolio, 2008; Uffelmann et al., 2021). Figure 1 schematically depicts how a typical GWAS unfolds. Researchers start by collecting DNA samples from both cases and controls. All DNA samples are genotyped using DNA microarrays and imputed through reference genomes to reconstruct

complete genotypes. Finally, researchers test for statistically significant differences between the genotypes of cases and controls. The genes harboring variants strongly associated with a disease can be suspected to play a role in its etiology, hence being potential targets for pharmaceutical intervention. Appendix B presents additional details and an example of a genome-wide association study.

Unlike candidate gene studies, where researchers decide which subset of genes to target, genome-wide association studies look for genetic variants across the whole genome (Visscher et al., 2017; Uffelmann et al., 2021). A genome-wide search permits to scan the entire set of possible gene-disease combinations, pointing directly to the most promising ones (Panel (a) of Figure 2). In practice, this search strategy removes one degree of freedom from the researcher, who is no longer required to specify genetic targets *ex-ante*. This ensures that GWASs are not biased by prior biological knowledge and beliefs, thus avoiding researchers' tendency to focus on familiar genes. Genome-wide association studies generate discoveries thanks to what directly emerges from the data, making them a prime example of data-driven search (Evans and Rzhetsky, 2010).

Genome-wide association studies have been harshly criticized for some shortcomings. On the one hand, these studies are inherently correlational, which means that any finding could be a false positive (Marigorta et al., 2018). On the other hand, even if the associations discovered by GWASs are statistically significant and replicable, scholars have suggested that this approach neglects more complex interaction structures between genes (Boyle et al., 2017). Moreover, most associations explain a small fraction of the genetic variation in disease susceptibility, which means that the therapeutic benefit from intervening in them could be quite small (Goldstein et al., 2009). These criticisms explain why candidate gene approaches remain popular among researchers, but it must be noted that the debate on whether GWASs discover scientifically impactful gene-disease associations is still unsettled.

4 Data

4.1 Information on Gene-Disease Associations

I construct a dataset of all novel gene-disease associations (GDAs) introduced from 1980 to 2016 inclusive. I retrieve such information from DisGeNET (v7.0), an aggregator considered a complete repository of scientific results linking human diseases to their genetic causes (Hermosilla and Lemus, 2019; Piñero et al., 2020). This database collects GDAs harvested from specialized sources, including curated datasets and publications indexed in PubMed. My data are at the GDA level, and for each association, I retrieve both the publication that introduced it and the list of

all follow-up articles that investigated it. I focus on associations mapping a protein-coding gene to a disease, syndrome, or abnormality with clear health implications. My final sample includes 349,670 gene-disease associations between 14,098 genes and 15,030 narrow disease categories.¹

To identify which associations are introduced with a data-driven approach, I rely on the GWAS Catalog, a manually curated source managed by the European Bioinformatics Institute (MacArthur et al., 2017). The GWAS Catalog is a comprehensive list of genome-wide association studies published in peer-reviewed journals. Studies are eligible for inclusion in the GWAS Catalog if they use a DNA microarray to scan the entire genome without targeting any specific gene *ex ante*. The Catalog also collects the details of the specific gene-disease associations tested in the study. Following the best research practices, only associations with a high statistical significance ($p\text{-value} < 1.0 \times 10^{-5}$) are considered (Marigorta et al., 2018). My sample includes 8,464 GDAs introduced by 1,223 distinct genome-wide association studies. Panel (b) of Figure 2 shows the rapid growth of GWASs since 2005, when the first such study was published.

Information on the bibliographic characteristics of papers introducing at least one novel gene-disease combination is taken from NIH’s iCite data. Specifically, I record the number of authors of each article, the journal, and the number of citations received. I use the 2020 SCImago journal ranking to measure the relative prestige of the publication venue of each paper. To identify the principal investigator (PI), I extract information on the last author of each publication from the Author-ity database (Torvik and Smalheiser, 2021).² Author-ity is a highly accurate database that disambiguates the authors of PubMed papers leveraging information on names, coauthors, MeSH codes, affiliations, and paper keywords. I use this data to estimate fixed effect models that identify the effect of carrying out a GWAS controlling for time-invariant characteristics of the principal investigator.

I also gather additional gene-level attributes. For each gene, I record if it is part of a gene family. Genes in a family are formed by duplication of a single ancestral gene and generally share similar biochemical properties (Daugherty et al., 2012). Such genes have the same name followed by a number reflecting the order in which they were discovered (e.g., BRCA1 and BRCA2, discovered in 1994 and 1995, respectively). Any discrepancy in attention between members of the same gene

¹Scientists routinely complain that associations proposed in academic publications often turn out not to be robust (Tabor et al., 2002). To avoid that this issue affects my analyses, I use the DisGeNET-provided *Evidence Index* to retain in my data only associations for which contradictory results represent less than 10% of the available publications about them. However, all my results are robust to either stricter thresholds of the Evidence Index or to keeping the whole DisGeNET data. See Appendix Figure C.4 for robustness checks.

²Authorship norms in the life sciences prescribe that the principal investigator is placed in last position on the authorship roster of a paper (Evans, 2010). The focus on principal investigators is justified by the fact that they have agency in directing the methodological choice of each study.

family is likely to reflect path dependence in studying the genes discovered earlier in time but not differences in scientific potential (Stoeger et al., 2018). This fact is supported by my data, where I observe that the first member of a gene family receives, on average, 56% more publications than the second member of the same family.

Next, I code which human genes have a homolog gene in the lab mouse (Clarke, 2002; Murray et al., 2016). Homologs are genes inherited in two species from a common ancestor, thus retaining comparable functions and biology. This property allows scientists to carry out experiments on homolog genes in animals to learn about human biology. Since the mouse is the most used scientific tool for gene knockouts (i.e., a lab technique to study the role of a gene by preventing its normal functioning), genes without a mouse homolog are less convenient to study experimentally and thus often neglected for reasons not related to their importance (Baba and Walsh, 2010; Stoeger et al., 2018). Finally, I code which genes regulate the function of more than one downstream gene (Türei et al., 2016). Studying the role of those genes in a disease requires learning how they influence several interdependent biological pathways, thus offering a potentially more complex route to therapeutic development (Hermosilla and Lemus, 2019).

4.2 Outcome Variables

My objective is to compare the characteristics of GWAS-established gene-disease associations vis-à-vis associations established with a candidate gene approach. To do so, I ask: conditionally on being introduced by a GWAS, do gene-disease associations present statistically different attributes? In particular, I focus on two outcomes:

Underexplored Gene: I use two alternative proxies to capture which genes received scant attention before the emergence of GWASs. The first dependent variable is a dummy that takes value one for gene-disease associations that include a gene never associated with a disease before 2005, the year of the first GWAS. The second proxy is the gene's discovery date, since many of the genes mapped by the Human Genome Project are still overlooked due to path-dependent research choices (Stoeger et al., 2018). Accordingly, I explore if GWASs are relatively more likely to introduce gene-disease combinations involving genes discovered after the year 2000 (when the first draft of the human genome was released). Both dependent variables are coded as dummies to allow a straightforward interpretation of the OLS coefficients as linear probability models, but I repeat the same analyses using continuous versions of these variables in the Appendix.

Scientific Impact: The second dependent variable is a dummy that takes value one for gene-disease associations that have a large scientific impact. Usually, researchers rely on paper-to-paper

citation counts to measure impact, but this would be inappropriate in my setting since scientific articles often study more than one gene-disease association. This prevents a straightforward way to assign to each GDA its share of the citations received by the focal article introducing it.³ Instead, I exploit DisGeNET to build a novel measure of GDA scientific impact: the number of papers that *directly* build on the gene-disease combination. These include empirical and experimental work that investigates the proposed association, regardless of whether they cite the paper that first introduced it, hence being a truthful measure of GDA impact. For each year, I code as high-impact all new GDAs in the 95th percentile of follow-on work received.⁴

4.3 Summary Statistics

Table 1 lists the key variables together with the summary statistics for the sample used in the analysis. Panel A provides summary statistics about the publications that established new GDAs in the period considered. Besides being more cited than candidate gene papers, on average GWASs introduce more associations spanning a larger number of genes. Panel B provides summary statistics at the GDA level. Previewing the following analysis, the incidence of high-impact associations is higher for GWASs than for candidate gene papers. It also appears that genes associated with a disease by GWASs are more likely to be less studied, to lack a mouse homolog, and be the second member of gene families. The share of breakthrough combinations that include an understudied gene is 13.8% among those established by candidate gene studies but grows to 37.6% among those found by genome-wide association studies.

5 Results

5.1 Data-Driven Search and the Direction of Innovation

I use OLS to estimate the following linear probability model using gene-disease level data: $\mathbb{I}(GDA \text{ with understudied gene} > 0)_i = \alpha + \beta \mathbb{I}(Introduced \text{ by GWAS} > 0)_i + \gamma \mathbf{X}_i + \epsilon_i$, where \mathbf{X}_i include disease and principal investigator fixed effect, as well as controls for year, journal prestige, and number of authors of the paper that introduced GDA i . $\mathbb{I}(GDA \text{ with understudied gene} > 0)_i$ is an indicator variable equal to one if GDA i includes an understudied gene. $\mathbb{I}(Introduced \text{ by GWAS} > 0)_i$ takes value one for GDAs introduced by a

³Moreover, citation counts would be misleading in this context since GWASs are highly cited on average ($\mu_{GWAS}=171$ vs. $\mu_{Candidate \text{ Gene}}=42$) due to a variety of reasons unrelated with the scientific quality of the findings, such as reviews, criticisms, or commentaries that discuss the results of the genome-wide approach.

⁴This choice follows the approach of papers on breakthrough innovation that operationalize outlier performance as falling within the top 5% of the sample (Arts and Fleming, 2018; Kaplan and Vakili, 2015). However, my results are unchanged if I use alternative cut-offs (Appendix Table C.2) or directly the count of follow-on papers as a dependent variable (Appendix Table C.3).

GWAS, and zero for GDAs introduced by candidate gene studies. This specification estimates the difference between GDAs that first appeared in a genome-wide association study and GDAs that were introduced by candidate gene papers. If data-driven search leads to diversify search, then I should find that the OLS estimate β is positive and statistically significant.⁵ All specifications cluster standard errors two-way at the gene and disease level.

Table 2 presents estimates from this regression. The main result is that genome-wide association studies are significantly more likely to associate understudied genes with human diseases than candidate gene studies. Specifically, the estimate of β in Column 1 indicates an average increase of 20 percentage points on the probability of combining a gene never associated with a disease before 2005, a significant increase given that the baseline is about 13 percentage points. Column 3 shows that this finding is robust to using the date of discovery as an alternative proxy for genes that have been historically less studied. However, a potential concern could be that scientists carrying out GWASs are systematically different from those who do not. If the characteristics of researchers that publish GWAS correlate with the likelihood of exploring less studied genes, then the results could reflect an upward bias. One way to ameliorate this concern is to include principal investigator fixed effects, thus controlling for time-invariant attributes that might confound the estimates. Columns 2 and 4 of Table 2 reassure that the results are robust to the addition of PI fixed effects: principal investigators that carry out a GWAS are 86-114% more likely to recombine understudied genes relative to when they adopt a candidate gene approach.⁶

Figure 3 presents an intuitive visualization of the combinatorial space of pairwise gene-disease combinations. Comparing the areas searched by candidate gene studies with the findings of genome-wide association studies illustrates the difference between the two strategies. New combinations introduced by GWASs span a much wider area of the technological landscape, while theory-driven search tends to replicate existing research patterns. Panel (b) also validates the global nature of GWAS: for each disease investigated, the range of genes associated spans the entire genome. However, the figure points to the fact that GWASs keep focusing on historically well-studied diseases (see also Figure C.1 in the Appendix). This result shows how data-collection decisions remain crucial in determining the direction of search, but also confirms that the diversification in gene space is due to the search strategy itself and not to a change in disease focus.

Focusing on a narrow subset of genes is not necessarily a problem if they are chosen because of

⁵The choice of using a linear probability model is motivated by the large number of fixed effects employed, which prevents convergence in nonlinear models.

⁶Appendix Table C.1 reports similar results when considering continuous versions of the dependent variables instead of a discrete coding. GWASs are more likely to recombine genes that received 31-34% less publications before 2005, and that were discovered 1.6-2.2 years later.

their higher scientific promise. To explore this possibility, I consider new associations recombining genes that are part of a gene family. Since genes in the same family are biologically similar, the main difference between the first and second family members is often just the order in which they were discovered (Stoeger et al., 2018). Figure 4 shows that candidate gene papers tend to study the first gene of a family much more than the second member. Such discrepancy reflects how scientists continue targeting the gene discovered earlier, even holding biological characteristics and therapeutic potential constant (see also Appendix Table C.4). However, the difference between the first and second genes in a family completely disappears for GDAs introduced by genome-wide association studies. This evidence suggests that one of the mechanisms through which GWASs help discovery is counteracting inertial forces in scientists’ research paths, which are especially damaging to our understanding of polygenic diseases (Haynes et al., 2018).

5.2 Does Data-Driven Search Lead to Better Innovations?

While data-driven search broadens the scope of search, its impact on the value of innovation is ambiguous. In this section, I estimate the following specification: $\mathbb{I}(GDA \text{ in top } 5\% \text{ of impact} > 0)_i = \alpha + \beta \mathbb{I}(Introduced \text{ by GWAS} > 0)_i + \gamma \mathbf{X}_i + \epsilon_i$, where $\mathbb{I}(GDA \text{ in top } 5\% \text{ of impact} > 0)_i$ is an indicator variable equal to one if GDA i is among the top 5% most impactful combinations, the common definition for breakthrough innovation. All other variables and controls are identical to the previous section’s specification.

Table 3 presents the results. In the basic specification reported in Column 1, new gene-disease associations that first appeared in a genome-wide association study are, on average, 24% more likely to be among those of high scientific impact. However, this result could just be a reflection of the change in genetic focus documented in the previous section. If the new genes recombined by GWASs are intrinsically more likely to yield high-value associations, then the increase in breakthrough innovation might mechanically derive from that. To investigate this possibility, I estimate the same model by adding gene fixed effects, hence absorbing the cross-sectional variation linked to genes’ scientific potential. Column 2 of Table 3 shows that the estimate’s magnitude and statistical significance substantially grow after adding gene fixed effects. Results are robust to the inclusion of PI fixed effects, ruling out that they are driven by time-invariant researchers’ attributes (Column 3 of Table 3).

In the Appendix, I further explore the sensibility of my findings. Table C.2 shows that results do not change if I adopt alternative ways to define breakthrough discoveries, such as gene-disease combinations in the top 10% or top 1% of scientific impact. If anything, the magnitude of the

coefficient grows larger for more stringent definitions of breakthrough, suggesting that GWASs are more powerful in uncovering outlier combinations. I also show that the results are unchanged if I use as a dependent variable the count of follow-on papers received by each combination instead of a dichotomous definition of breakthroughs (Table C.3). Finally, I find the same results if I limit my analysis to GDAs involving members of gene families (Table C.4). In sum, my analysis documents that gene-disease associations introduced by GWASs have on average a higher scientific impact than comparable associations discovered with candidate gene approaches, even when holding the intrinsic characteristics of the genes and the attributes of the principal investigator constant.

5.3 Data, Theory, and Landscape Characteristics

The results reported in the previous sections capture the average effect of GWAS on discovery, but this could conceal large heterogeneity in the effectiveness of data-driven search. In particular, one might expect that theory-driven search becomes more effective in areas where scientists have more profound biological knowledge. In those domains, theoretical knowledge should lead researchers directly to the best technological combinations and outperform data-driven search (Arora and Gambardella, 1994; Fleming and Sorenson, 2004). Figure 5 explores this idea by plotting the likelihood of introducing high-impact GDAs in correspondence with each gene, separately by the search strategy that introduced them. Genes on the X-axis are sorted by the number of pre-2005 publications received, which can serve as a proxy for the depth of biological knowledge available. A striking pattern emerges: while the ability of GWASs to introduce valuable gene-disease combinations is roughly constant across the genetic landscape, candidate gene studies are much more effective for genes that have received more study in the past.⁷

A potential drawback of this analysis is that the distribution of theoretical expertise on the landscape is endogenous (Gittelman, 2016; Nelson, 2003). Genes more heavily studied are likely to be those considered most relevant by scientists, hence this might explain why theory-driven search performs better in that case. To rule out this concern, I exploit the fact that some genes have historically been neglected simply because they cannot be studied with the lab mouse (Stoeger et al., 2018). This simple intuition is borne by my data, because I find that genes without a counterpart in the mouse genome have received, on average, 22% less publications on how they might be implicated in human diseases. Since these genes are less theoretically characterized due to only to experimental convenience, this provides variation in theory availability that is unrelated to their therapeutic potential. Figure 6 shows that, unlike GWAS, candidate gene studies are markedly less successful

⁷The same pattern emerges if I use the date of discovery of the gene as a proxy for the availability of biological theory, see Appendix Figure C.2.

when exploring genes less theoretically known because they lack a mouse homolog. This pattern is confirmed by linear probability models that control for disease, gene, and principal investigator fixed effects (Table 4).

The analyses reported above established that theory-driven search is more effective in areas where scientists have more profound theoretical knowledge, while data-driven search is powerful in recombining little-known components. But what is the precise mechanism behind this finding? One hypothesis is that there are landscape characteristics that moderate the relative effectiveness of the two search strategies, such as the density of interrelationships between components (Gavetti and Levinthal, 2000; Fleming and Sorenson, 2001; Kapoor and Wilde, 2022). I explore this possibility by estimating the efficacy of genome-wide association studies when they involve genes that regulate the function of more than one other gene. Intuitively, when a gene governs the function of many downstream genes, it becomes complex to figure out the precise mechanism through which it is related to a disease (Hermosilla and Lemus, 2019). The results in Table 5 highlight that the relative advantage of candidate gene studies in the context of highly-studied genes is exclusively due to their ability to recombine genes involved in many regulatory networks. This suggests a key limitation of data-driven search: data signals report correlations that are unable to discern complex interactions between interdependent or coupled components (Fleming and Sorenson, 2004; Ghosh, 2021). Instead, in the presence of a solid-enough theoretical understanding, researchers can locate the best gene-disease combinations because they can account for how genes interact (Boyle et al., 2017).

5.4 Robustness Checks

A. Characteristics of Scientists Publishing GWASs: The inclusion of principal investigator fixed effects allows estimating within-researchers effects, capturing the change in outcomes experienced by the same researcher after switching search strategy. However, it could still be that researchers who are systematically more explorative or able to identify breakthroughs sort into adopting GWASs, leading to a potential upward bias in the estimates. Appendix Figure C.3 shows that before their first GWAS, principal investigators that will eventually adopt a genome-wide search are not more likely to target less studied genes or to introduce breakthrough gene-disease combinations. This robustness test ameliorates concerns that researchers sort into carrying out a GWAS based on endogenous characteristics related to my outcomes of interest.

B. Disease Selection Over Time: One additional concern is that my analyses could overstate the benefits of adopting GWAS if the search method is applied to diseases where it is more likely to

yield discoveries involving unknown genes. Note that Figures 3 and C.1 already show that GWAS target the same highly-studied diseases as the majority of candidate gene studies, hinting that this might not be an issue in practice. Appendix Table C.5 directly tests this idea, showing that diseases that receive their first GWAS earlier in my sample period are no more likely to be associated with short-changed genes. Looking at the list of diseases that received the most GWASs, one finds polygenic diseases with a large incidence in the population, suggesting that this was the primary criterion guiding disease choice.

C. Alternative Samples: Gene-disease associations require extensive follow-on work to be validated and contradictory results on their robustness are not infrequent (Uffelmann et al., 2021). The sample used in this paper considered associations for which DisGeNET reports less than 10% of contradictory papers about them, to avoid confounding effects from false positive discoveries. In Figure C.4 I test the robustness of the main results to different selections of the sample, ranging from all DisGeNET associations to the inclusion of only those for which no contrasting evidence exists. Results are quantitatively similar regardless of the sample chosen.

D. Different Measure of Scientific Value: Instead of relying on the number of subsequent publications to measure associations' scientific potential, I perform a robustness check using DisGeNET's *GDA Score* (Piñero et al., 2020). The GDA Score synthetically captures the scientific reliability of all the existing evidence on the gene-disease association. Table C.6 presents the coefficient of the OLS regressions for each of the subsamples of genes analyzed in Section 5.2. The results confirm the earlier findings on the effectiveness of data-driven search in introducing combinations of higher scientific value.

E. Therapeutic Value of GWAS Findings: The analyses reported so far showed that GWAS uncover GDAs with a higher scientific value on average, but this does not guarantee that those gene-disease pairs will also prove therapeutically useful. To capture downstream medical translation, I code a dummy that captures whether papers introducing new gene-disease combinations are later cited by articles reporting clinical trials. Appendix Table C.7 reports that genome-wide association studies are more likely to be cited by clinical trial articles than comparable candidate gene studies. This finding reassures that findings scientifically impactful are also those with larger value to developing treatments.

6 Conclusion

In this paper, I explore how data changes the search for innovation. Unlike theory-driven search, I argue that data enables global search strategies that lead to more exploratory recombinations of technological elements. Empirical results in the context of genome-wide association studies confirm this idea, showing that path dependency does not tether data-driven search. Data lead innovators to experiment with short-changed areas of the technological landscapes and help them uncover combinations of higher average value. The latter result is stronger in uncharted landscape areas, but theory-driven approaches are more effective when deeper theoretical knowledge can be used to guide search. The key mechanism underlying this finding is the ability of theory to account for complex interdependencies among technological components. Instead, data signals report misleading correlations when dealing with highly coupled components.

My work makes several contributions. First, I provide a theoretical and empirical framework to explore data-driven innovation. Data can drastically change the “technology of technical change” (Arora and Gambardella, 1994; Agrawal et al., 2022; Cockburn et al., 2019), but to date, there is little understanding of how data analytics is reshaping the generation of novelty (Allen, 2022; Cao et al., 2021; Hoelzemann et al., 2022; Nagaraj, 2022; Wu et al., 2020). Addressing this gap is of first-order importance in the age of big data, and my work is a first step in this direction. Second, the construct of data-driven search constitutes an addition to the theory of recombinant search (March, 1991; Fleming and Sorenson, 2001; Gavetti and Levinthal, 2000; Katila and Ahuja, 2002; Levinthal, 1997; Schilling and Green, 2011). By framing data as reshaping recombinant search, I leverage this important body of research to clarify the conditions under which theory-driven search will outperform data-driven search. Moreover, my study is one of the first to empirically measure how alternative search strategies map into discoveries (Kneeland et al., 2020; Maggitti et al., 2013). Finally, I provide new empirical evidence on how data analytics in genomics are reshaping pharmaceutical innovation (Gambardella, 1995; Hermosilla and Lemus, 2019; Kao, 2022). My results are timely also given the lack of consensus about the scientific value of genome-wide association studies (Boyle et al., 2017; Pearson and Manolio, 2008; Visscher et al., 2017).

This paper has practical implications for scientists, managers, and governments. For individual researchers, my results highlight when alternative search strategies are more or less effective, suggesting that data analytics should be relied upon, especially when venturing into uncharted domains. More in general, data analytics are diffusing in every sector of the economy, but the returns remain heterogeneous and concentrated among few companies (Brynjolfsson et al., 2021). This paper highlights that managers should collect and rely on data to guide risky experimentation

but also trust their knowledge base when working in well-trodden domains. My results also provide an additional rationale for furthering investments in large-scale public data sources that might enable data-driven search (Nagaraj, 2022; Kao, 2022).

It is also important to note that despite its considerable potential, data-driven search is not a panacea for recombinant search. Exclusive reliance on data could end up hindering search, either because it can lead scholars to look only “where the light is” (Hoelzemann et al., 2022) or because blind reliance on available data might replicate their biases (Cao et al., 2021). Moreover, data might be of little help when trying to invent entirely new technologies that do not emerge from old components (Wu et al., 2020). As such, data-driven search constitutes a within-paradigm search that might be subject to technological exhaustion unless new elements are added over time (Fleming, 2001).

Finally, despite the contributions outlined above, a few limitations of this paper must be acknowledged. First, the present study allows for meaningful comparisons of the consequences of engaging in theory-driven versus data-driven search, but the research design does not fully account for the endogenous choice of the form of search. Second, the documented patterns are from the quantitative case study of a single domain. Locating the genetic roots of human diseases is crucial for drug discovery, but it has specificities that might not directly translate to other settings. While an increasing number of domains are receiving complete maps of the relevant technological landscapes, just like the Human Genome Project did for the genome, data-driven search might remain unfeasible in other contexts. More research will be needed to investigate the external validity of my findings. Finally, my work does not explore how data-driven findings affect downstream investments in new drugs. This is an exciting avenue for follow-up work that is outside the scope of this paper.

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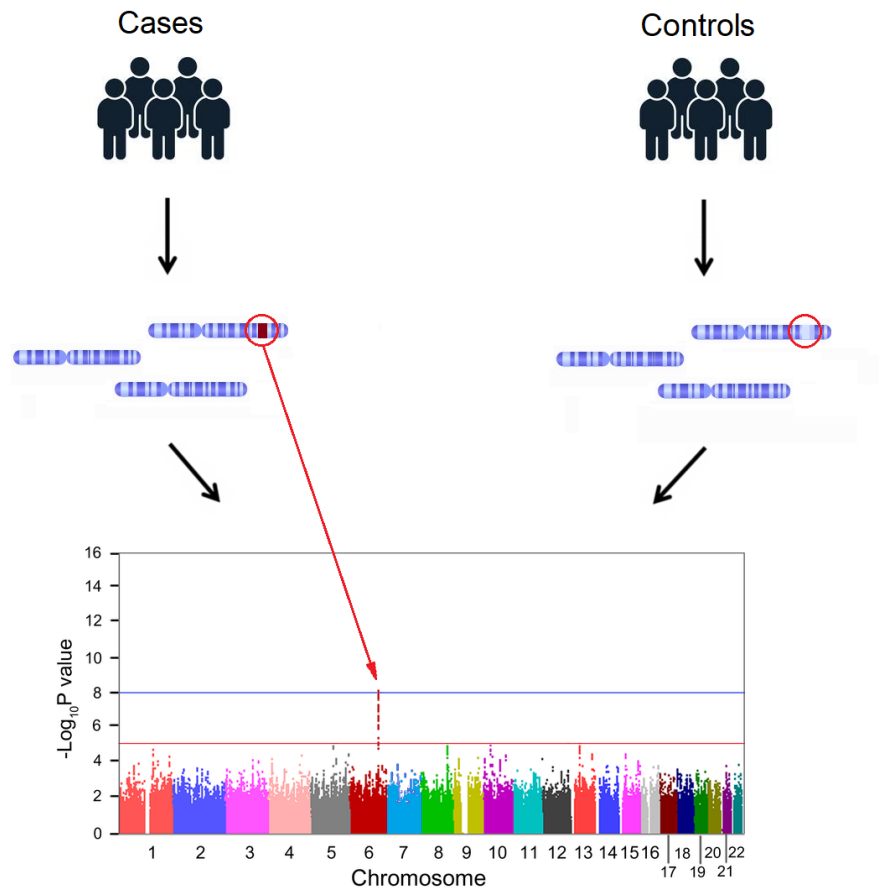
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7 Figures and Tables

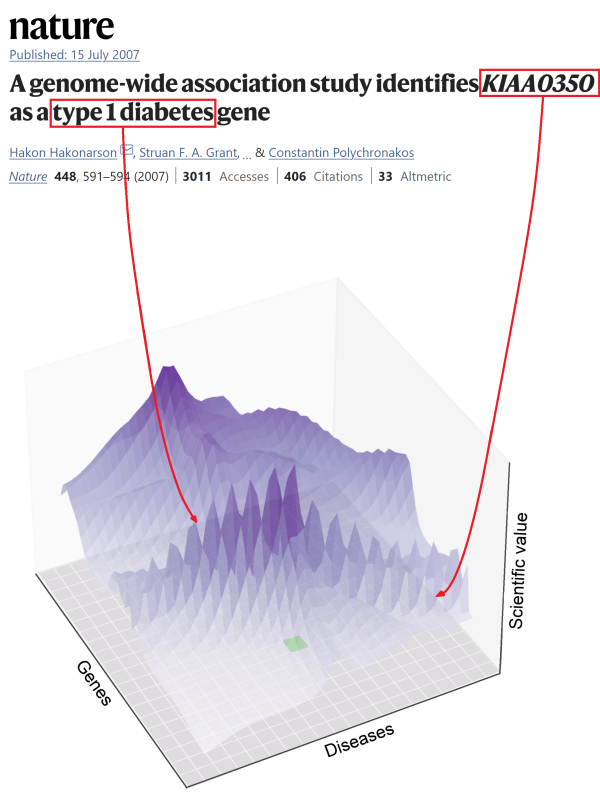
Figure 1: Schema of how a typical genome-wide association study unfolds.



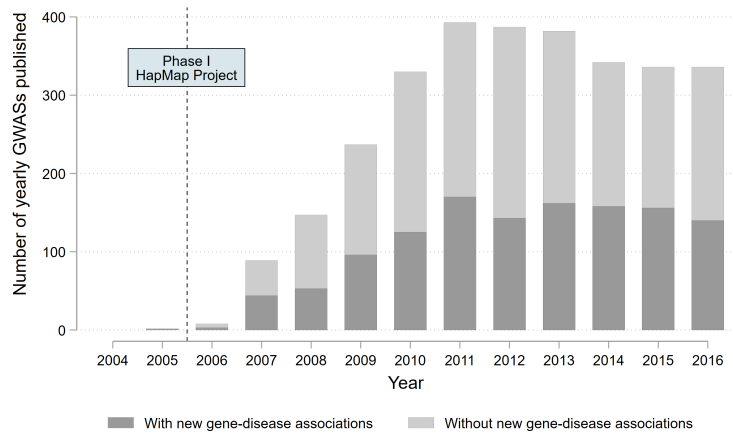
Note: The figure depicts the main steps involved in a genome-wide association study. First, the researchers select the disease of interest and assemble a group of cases (subjects showing the condition) and one of controls (healthy subjects). Then, the genome of people with and without the condition are sequenced in search of significant genetic differences. Finally, statistical methods are used to test the association between any genetic variant and the disease of interest. The panel at the bottom is the characteristic “Manhattan plot” which indicates the location of the statistically significant genetic variants in the chromosome. On the Y axis there is the strength of the finding expressed as $-\log_{10}(\text{p-value})$, hence higher values correspond to stronger associations. For instance, the picture depicts a genetic variant in chromosome 6 that is significantly associated with the condition.

Figure 2: The emergence of genome-wide association studies in the search for useful gene-disease combinations.

(a) GWAS as data-driven search

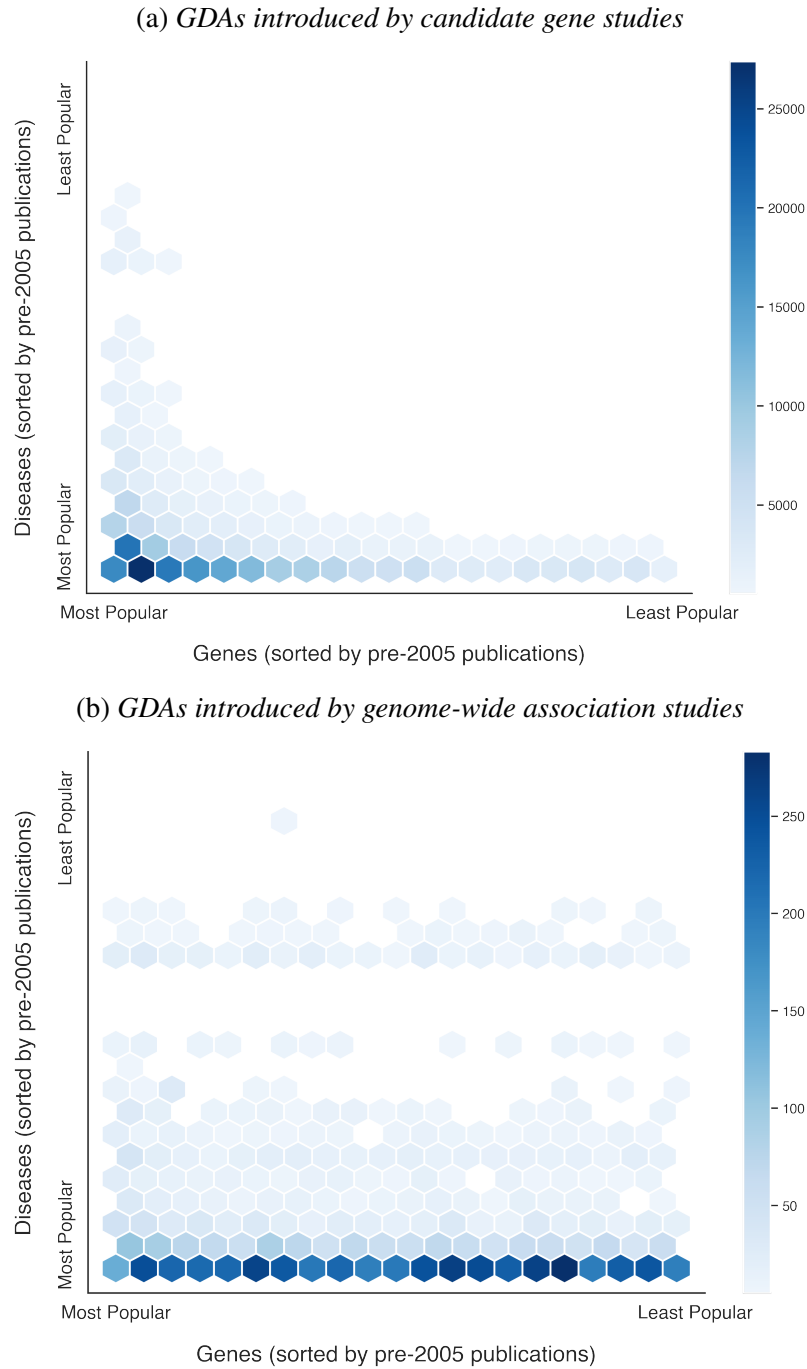


(b) Yearly GWASs published



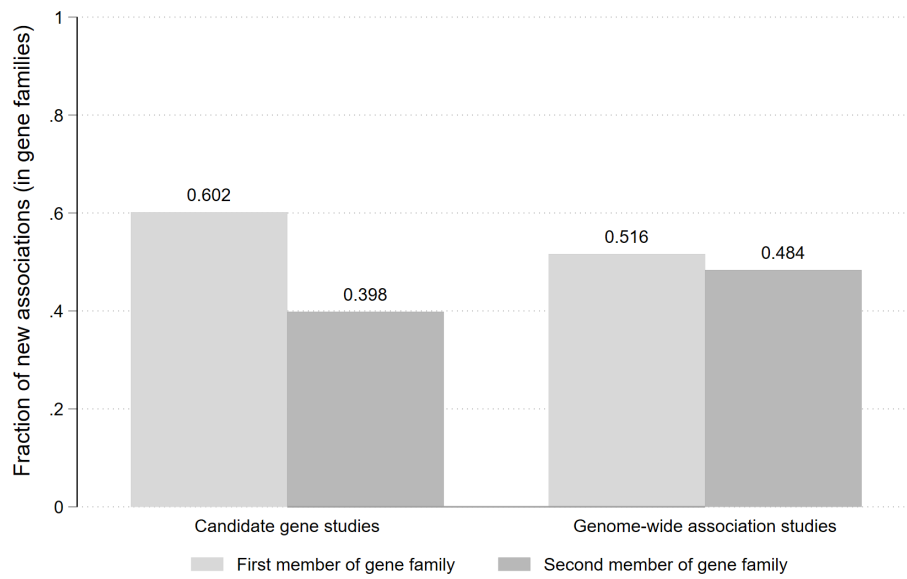
Note: Panel (a) depicts how a typical GWAS introduces a new gene-disease associations in the combinatorial landscape. Each combination of gene and disease has a specific scientific value, captured by the elevation at that location. Details on the GWAS by Hakonarson et al. (2007) are available in the Appendix B.3. Panel (b) shows the number of yearly genome-wide association studies published. The dark grey portion of the bars is the number of GWASs that introduced at least one novel gene-disease association, thus being the object of the present study. Data are from the GWAS Catalog. The vertical dashed line marks the completion of the Phase I of the HapMap project in 2005. See text for details.

Figure 3: Genome-wide associations studies span a larger portion of the genetic landscape relative to candidate gene studies.



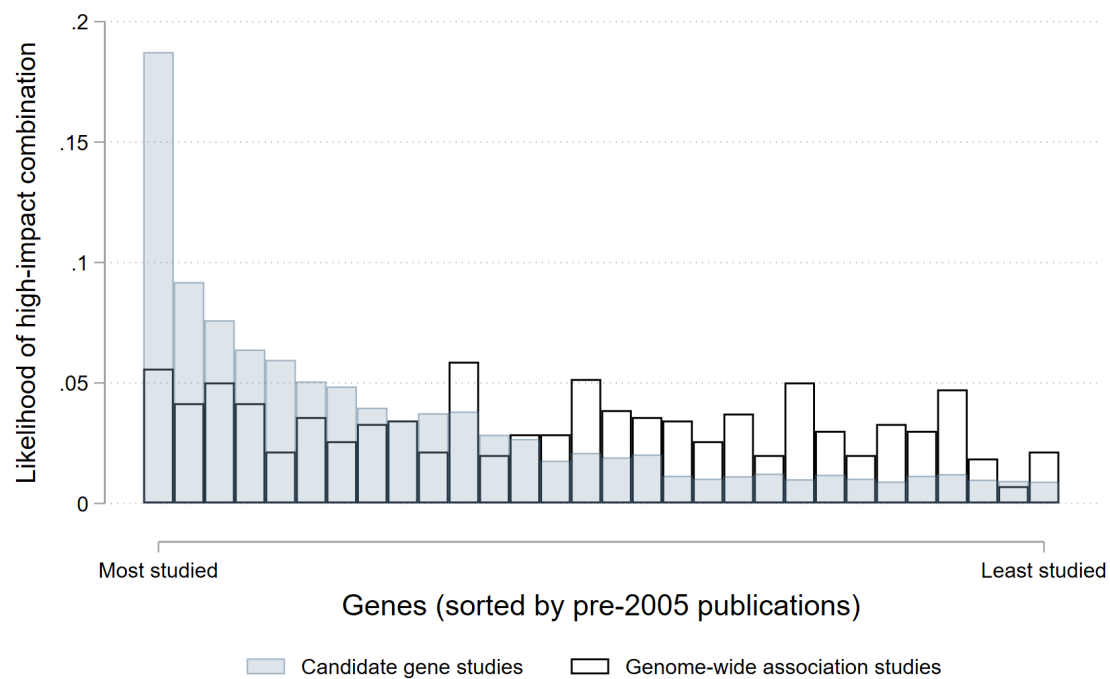
Note: Panel (a) shows a heatmap of new gene-disease associations introduced after 2005 with candidate gene studies. Panel (b) shows a heatmap of new gene-disease associations introduced after 2005 with genome-wide association studies. Both panels have 14,112 genes on the X axis, sorted from the most to the least studied in the pre-GWAS era, and 15,039 disease categories on the Y axis, sorted from the most to the least studied in the pre-GWAS era. Darker areas denote the introduction of a higher number of new gene-disease associations. See text for details.

Figure 4: Gene-disease associations introduced by a GWAS are not biased towards the first member of gene families.



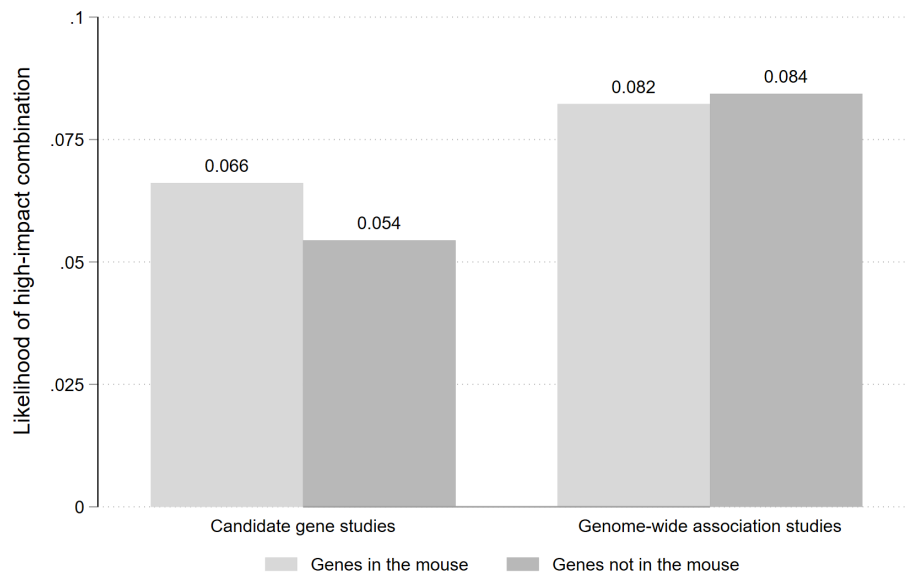
Note: The figure plots the share of new gene-disease associations on the first vs. the second member of a gene family, conditional on involving a gene family member. Data used in the graph are limited to all new gene-disease associations introduced in the period 2005-2016. Only diseases targeted by at least one GWAS are considered in this figure. See text for details.

Figure 5: Gene-disease associations introduced by GWAS are more likely to be high-impact for understudied genes, but candidate gene studies perform better when genes are well studied.



Note: The histogram plots the share of high-impact GDAs for each gene distinguishing by the type of study that introduced them. Data used in the graph are limited to all new gene-disease associations introduced in the period 2005-2016. The 14,112 genes on the X axis are sorted from the most to the least studied in the pre-GWAS era. See text for details.

Figure 6: GWAS are especially effective to introduce high-impact gene-disease associations for genes with less theoretical knowledge because they cannot be studied with the lab mouse.



Note: The figure plots the share of high-impact GDAs for genes with and without a mouse homolog distinguishing by the type of study that introduced them. Genes that lack a homolog gene in the lab mouse receive less attention and are more likely to be little known, but not because they are less biologically important. Data used in the graph are limited to all new gene-disease associations introduced in the period 2005-2016. Only diseases targeted by at least one GWAS are considered in this figure. See text for details.

Table 1: Descriptive statistics.

| Panel A: paper-level descriptives | | | | | | | | | | | | |
|-----------------------------------|------------------------|--------|--------|------|-------|---------|---------|--------|---------|------|-------|-------|
| | Candidate gene studies | | | | | | GWAS | | | | | |
| | mean | median | st d | min | max | N | mean | median | st d | min | max | N |
| Forward citations | 41.50 | 22 | 84.389 | 0 | 8,084 | 136,550 | 171.21 | 81 | 292.771 | 0 | 2,822 | 1,223 |
| Rank of the journal (ventile) | 13.25 | 14 | 5.061 | 1 | 20 | 136,550 | 17.52 | 19 | 3.763 | 2 | 20 | 1,223 |
| Associations per paper | 4.78 | 4 | 6.924 | 1 | 927 | 136,550 | 10.48 | 5 | 20.509 | 1 | 307 | 1,223 |
| Genes per paper | 2.15 | 2 | 2.150 | 1 | 690 | 136,550 | 6.74 | 3 | 12.024 | 1 | 169 | 1,223 |
| Number of authors | 9.01 | 8 | 6.549 | 1 | 445 | 136,550 | 38.67 | 25 | 44.762 | 1 | 565 | 1,223 |
| Year | 2011.10 | 2011 | 3.392 | 2005 | 2016 | 136,550 | 2012.32 | 2012 | 2.506 | 2005 | 2016 | 1,223 |

| Panel B: association-level descriptives | | | | | | | | | | | | |
|--------------------------------------------------|------------------------|--------|-------|------|------|---------|---------|--------|-------|------|------|-------|
| | Candidate gene studies | | | | | | GWAS | | | | | |
| | mean | median | st d | min | max | N | mean | median | st d | min | max | N |
| With never recombined genes (%) | 0.123 | 0 | 0.328 | 0 | 1 | 349,670 | 0.406 | 0 | 0.491 | 0 | 1 | 8,464 |
| With recently discovered gene (%) | 0.099 | 0 | 0.299 | 0 | 1 | 349,670 | 0.258 | 0 | 0.438 | 0 | 1 | 8,464 |
| With 2 nd member of a gene family (%) | 0.391 | 0 | 0.488 | 0 | 1 | 88,962 | 0.484 | 0 | 0.499 | 0 | 1 | 2,117 |
| With genes lacking mouse homolog (%) | 0.051 | 0 | 0.220 | 0 | 1 | 343,387 | 0.060 | 0 | 0.237 | 0 | 1 | 8,328 |
| In top 5% of scientific impact (%) | 0.049 | 0 | 0.216 | 0 | 1 | 349,670 | 0.083 | 0 | 0.276 | 0 | 1 | 8,464 |
| With regulator genes (%) | 0.501 | 0 | 0.500 | 0 | 1 | 349,670 | 0.0235 | 0 | 0.424 | 0 | 1 | 8,464 |
| Year of the association | 2011.08 | 2011 | 3.382 | 2005 | 2016 | 349,670 | 2012.68 | 2013 | 2.602 | 2005 | 2016 | 8,464 |

Note: Panel A presents descriptive statistics on papers that introduce new gene-disease associations after 2005. *Forward citations*= citations received by the focal article up to 2020 inclusive (data from NIH iCite); *Rank of the journal*= ventile of journal prestige (data from SCImago Journal Rank); *Associations per paper*= number of new GDAs introduced by the article; *Genes per paper*= number of genes associated with a disease by the article; *Number of authors*= number of authors co-authoring the article. Panel B presents descriptive statistics on new gene-disease associations introduced after 2005. *With never recombined genes (%)*= share of GDAs that include a gene never associated with a disease before 2005; *With recently discovered genes (%)*= share of GDAs that include a gene discovered after the year 2000 (i.e., after the Human Genome Project); *With 2nd member of a gene family (%)*= share of GDAs that include the second member of a gene family, conditional on being about a gene family (data on gene families from Stoeger et al. 2018); *With genes lacking mouse homolog (%)*= share of GDAs that include a gene lacking a gene homolog in the mouse (data from NIH); *In top 5% of scientific impact (%)*= share of GDAs that fall in the top 95th percentile of follow-on work (by year of discovery); *With regulator genes (%)*= share of GDAs that include a gene involved in controlling the function of more than one other gene (data from Türei et al. 2016); *Year of the association*= year in which the article introducing the GDA is published. See text for details.

Table 2: Genome-wide association studies are more likely to introduce gene-disease associations involving less-studied genes than candidate gene studies.

| Dependent Variable: | I(GDA for never associated gene>0) | | I(GDA for recently discovered gene>0) | |
|---------------------------|------------------------------------|-----------------------|---------------------------------------|-----------------------|
| GWAS | 0.200*** (0.01399) | 0.148*** (0.01563) | 0.112*** (0.01108) | 0.089*** (0.01406) |
| Disease FE | YES | YES | YES | YES |
| Principal Investigator FE | NO | YES | NO | YES |
| Journal prestige FE | YES | YES | YES | YES |
| Year of discovery FE | YES | YES | YES | YES |
| Number of authors FE | YES | YES | YES | YES |
| N | 352,162 | 331,825 | 352,162 | 331,825 |
| Mean of the DV: | 0.130 | 0.130 | 0.103 | 0.103 |
| Number of diseases: | 9,740 | 9,362 | 9,740 | 9,362 |
| Number of genes: | 14,072 | 13,910 | 14,072 | 13,910 |

Note: *, **, *** denote significance at 10%, 5% and 1% level respectively. Observations at the gene-disease association (GDA) level. Std. err. clustered two-way at the gene and disease level. $I(GDA \text{ for never associated gene} > 0): 0/1 = 1$ if the gene-disease association involves a gene never associated with a disease before 2005; $I(GDA \text{ for recently discovered gene} > 0): 0/1 = 1$ if the gene-disease association involves a gene discovered after the year 2000; $GWAS: 0/1 = 1$ for GDAs introduced by a genome-wide association study. See text for details.

Table 3: Genome-wide association studies are more likely to introduce gene-disease associations of high scientific impact than candidate gene studies.

| Dependent Variable: | I(GDA in the top 5% of scientific impact > 0) | | |
|---------------------------|--------------------------------------------------|-----------------------|-----------------------|
| GWAS | 0.012* (0.00663) | 0.037*** (0.00719) | 0.044*** (0.00918) |
| Disease FE | YES | YES | YES |
| Gene FE | NO | YES | YES |
| Principal Investigator FE | NO | NO | YES |
| Journal prestige FE | YES | YES | YES |
| Year of discovery FE | YES | YES | YES |
| Number of authors FE | YES | YES | YES |
| N | 352,162 | 350,693 | 330,250 |
| Mean of the DV: | 0.05 | 0.05 | 0.05 |
| Number of diseases: | 9,740 | 9,726 | 9,347 |
| Number of genes: | 14,072 | 12,617 | 12,463 |

Note: *, **,*** denote significance at 10%, 5% and 1% level respectively. Observations at the gene-disease association (GDA) level. Std. err. clustered two-way at the gene and disease level. $I(\text{GDA in the top 5\% of scientific impact} > 0):0/1=1$ if the gene-disease association involves a gene in the top 95th percentile of follow-on work (by year of discovery); $\text{GWAS}:0/1=1$ for GDAs introduced by a genome-wide association study. See text for details.

Table 4: Genome-wide association studies are especially more likely to introduce gene-disease associations of high scientific impact for genes less studied because they lack a mouse homolog.

| Dependent Variable: | I(GDA in the top 5% of scientific impact>0) | | | |
|---------------------------|---------------------------------------------|-----------------------|------------------------|-----------------------|
| Subsample: | Genes in the mouse | | Genes not in the mouse | |
| GWAS | 0.036*** (0.00738) | 0.044*** (0.00942) | 0.062* (0.03064) | 0.236*** (0.07017) |
| Disease FE | YES | YES | YES | YES |
| Gene FE | YES | YES | YES | YES |
| Principal Investigator FE | NO | YES | NO | YES |
| Journal prestige FE | YES | YES | YES | YES |
| Year of discovery FE | YES | YES | YES | YES |
| Number of authors FE | YES | YES | YES | YES |
| N | 326,506 | 306,232 | 15,848 | 11,478 |
| Mean of the DV: | 0.05 | 0.05 | 0.04 | 0.04 |
| Number of diseases: | 9,480 | 9,069 | 1,952 | 1,467 |
| Number of genes: | 11,502 | 11,361 | 743 | 695 |

Note: *, **, *** denote significance at 10%, 5% and 1% level respectively. Observations at the gene-disease association (GDA) level. Std. err. clustered two-way at the gene and disease level. $I(GDA \text{ in the top } 5\% \text{ of scientific impact} > 0): 0/1=1$ if the gene-disease association involves a gene in the top 95th percentile of follow-on work (by year of discovery); $GWAS: 0/1=1$ for GDAs introduced by a genome-wide association study. The sample used in columns 1 and 2 includes only genes with a homolog gene in the lab mouse, while the sample used in columns 3 and 4 includes only genes without a homolog gene in the lab mouse. See text for details.

Table 5: Genome-wide association studies are less likely to introduce high-impact gene-disease associations than candidate gene studies for genes involved in complex regulatory networks, provided that there is enough theoretical knowledge about them.

| Dependent Variable: | I(GDA in the top 5% of scientific impact>0) | | | | | |
|--------------------------|---------------------------------------------|-----------------------|-------------------------------------|-----------------------|-------------------------------------|-----------------------|
| Subsample: | Genes in top 50% of most studied | | Genes in top 75% of most studied | | Genes in top 90% of most studied | |
| GWAS | 0.038*** (0.01172) | 0.047*** (0.01590) | 0.031 (0.01987) | 0.072** (0.03251) | -0.032* (0.01750) | -0.001 (0.02411) |
| GWAS × Regulator Gene | | -0.016 (0.02071) | | -0.072** (0.03213) | | -0.057** (0.02501) |
| Disease FE | YES | YES | YES | YES | YES | YES |
| Gene FE | YES | YES | YES | YES | YES | YES |
| Journal prestige FE | YES | YES | YES | YES | YES | YES |
| Year of discovery FE | YES | YES | YES | YES | YES | YES |
| Number of authors FE | YES | YES | YES | YES | YES | YES |
| N | 176,206 | 176,206 | 86,421 | 86,421 | 33,159 | 33,159 |
| Number of diseases: | 7,901 | 7,901 | 6,298 | 6,298 | 4,463 | 4,463 |
| Number of genes: | 2,189 | 2,189 | 642 | 642 | 175 | 175 |

Note: *, **, *** denote significance at 10%, 5% and 1% level respectively. Observations at the gene-disease association (GDA) level. Std. err. clustered two-way at the gene and disease level. $I(GDA \text{ in the top } 5\% \text{ of scientific impact} > 0): 0/1=1$ if the gene-disease association involves a gene in the top 95th percentile of follow-on work (by year of discovery); $GWAS: 0/1=1$ for GDAs introduced by a genome-wide association study; $Regulator \text{ Gene}: 0/1=1$ for genes that are involved in controlling the function of more than one other gene (data from Türei et al. 2016). Column 1 and 2 include only GDAs with genes that received an above median number of studies before 2005. Column 3 and 4 include only GDAs with genes that are in the top 75% of the distribution of studies received before 2005. Column 5 and 6 include only GDAs with genes that are in the top 90% of the distribution of studies received before 2005.

Data-Driven Search and Innovation

Appendix

Matteo Tranchero

UC Berkeley

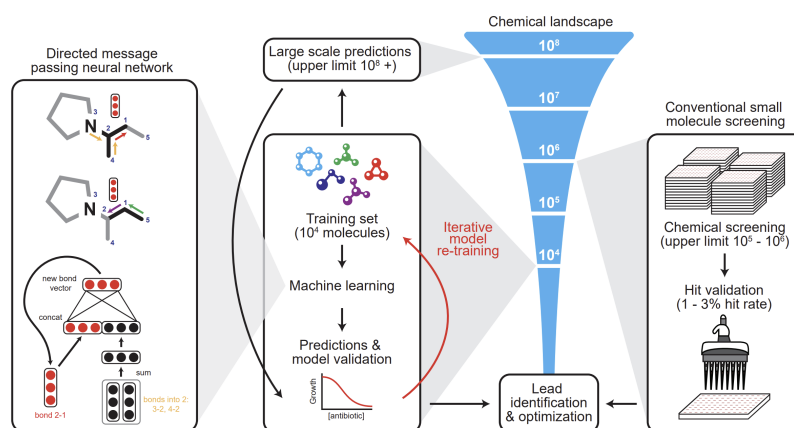
A An Example of Data-Driven Search from Combinatorial Chemistry

Consider the critical task of discovering new drugs. This is a difficult problem since the chemical space is complex and high-dimensional (Gambardella, 1995; Jayaraj and Gittelman, 2018; Rzhetsky et al., 2015). Historically, most successful molecules were serendipitously identified with random search or by experimenting in the neighborhood of known ones. This process was very long, costly, and inefficient (Arora and Gambardella, 1994; Gittelman, 2016). More recently, high-throughput screening (HTS) of large synthetic chemical libraries has opened up the possibility of large-scale rapid testing of millions of molecules. This approach usually involves pre-selecting a subset of the chemical space with drug-like characteristics (e.g., small molecules with a weight theoretically deemed to be suitable for human-use drugs) and sequentially testing them in physical assay plates (Jayaraj and Gittelman, 2018). Yet, records of this approach have been mixed since libraries are costly to maintain and the sequential screening has proved hard to scale to larger chemical spaces (Le Fanu, 2011).

However, new computational approaches and databases might provide an alternative for *in silico* testing that is not constrained by physical capacity or the compound libraries available. In a recent paper, Stokes et al. (2020) used a neural network approach to find molecules with antibacterial activity. Using data on known molecules to predict the bactericidal properties of structurally divergent molecules, the study’s authors discovered a new compound called halicin that has promising antibiotic properties. This result was achieved with a fraction of the time and costs involved in sequential assay screening, since it happened *in silico* without the need to undertake costly experimentation in the lab. Furthermore, this achievement is all the more remarkable considering that until then, no clinical antibiotics had been discovered using targeted high-throughput screening (Stokes et al., 2020).

Besides being a consequential example, the discovery of halicin also highlights the conditions

Figure A.1: Figure schematically representing the analysis of Stokes et al. (2020).



Note: The figure shows Figure 1 from Stokes et al. (2020). Using a deep neural network model, Stokes et al. (2020) built a molecular representation based on a specific property, in this case the inhibition of the growth of *E. coli*. Then, they applied the model to multiple chemical libraries to identify potential lead compounds with activity against *E. coli*. Finally, Stokes et al. (2020) selected a list of promising candidates after ranking the candidates according to the score predicted by the model.

under which data-driven search is feasible. First, the relevant characteristics of all the potential components to recombine must be observable. In the case of drug discovery, for example, this translates into the need to measure the structural properties of chemical compounds screened. While almost tautological, this first condition restricts the scope of data-driven search to settings in which components are identifiable and measurable *ex ante*. Second, there has to exist an agreed-upon metric of technological potential on which the promise of each combination can be assessed. This constitutes the objective function that data-driven search tries to maximize by finding the candidate combinations that score highest on it. For Stokes et al. (2020) this was the growth inhibition of *Escherichia coli*, but it is worth noting that a search guided only by data might not be feasible in fuzzier contexts where even the outcomes of the problem are ill-defined. Third, it must be possible to foresee the effect of novel combinations on the objective of interest. Going back to the example of antibiotics, this pertained to the prediction of whether a new compound could inhibit the growth of bacteria based on its structure.

B Genome-Wide Association Studies: Examples and Details

B.1 Genetic Research Before GWAS

Genomics is the branch of biological sciences concerned with the study of genomes, i.e., the entire collection of an organism's genes. In turn, genes are sequences of DNA bases that encode the "instructions" to synthesize gene products (e.g., proteins). Genes have a fundamental role in

the functioning of the human body, but sometimes they can acquire mutations in their sequence. When this happens, genes might alter their behavior and affect phenotypic traits, sometimes with significant consequences and the emergence of severe health conditions. However, the role of genes in the etiology of diseases offers an avenue for therapeutic intervention since genes associated with a condition can often be used as drug targets (Nelson et al., 2015). When a drug molecule binds to the therapeutic target it can modify its functioning, favorably affecting the outcome of a disease. Therefore, knowing the genetic roots of diseases has important practical consequences in the design of pharmaceutical drugs.

Genetic research has historically been very effective in locating genes individually responsible for disease conditions. These are called *Mendelian disorders*, which can usually be seen since birth and be deduced based on family history. Consider, for instance, the case of cystic fibrosis. This rare disorder can be caused by multiple DNA mutations which however cluster in a well-defined area of the genome, namely in the *Cystic fibrosis transmembrane conductance regulator* (CFTR) gene. Studying this kind of Mendelian disorders was one of the earliest and more significant successes of family studies in genetics (Bush and Moore, 2012). The primary research strategy involved genotyping families affected by cystic fibrosis: even with very small sample sizes due to the rarity of the disease, the very strong effect of the CFTR gene mutations allowed to identify the gene as causally connected with the disease unambiguously.

However, Mendelian diseases are typically rare because they tend to be eliminated by evolutionary pressures due to their gravity. Much more common are *complex diseases* that are not due to a single genetic factor, but rather by many genes. For complex diseases, any genetic mutation can increase the risk even without being neither necessary nor sufficient, which means that it is responsible for only a small proportion of the heritability of the disease. Although complex disorders often cluster in families too, they do not have a predictable inheritance pattern since they are influenced by convoluted interactions between genes and environmental factors. For this reason, family linkage studies have not fared well when applied to more common disorders.

B.2 A Scientific Primer on GWAS

To make progress in the study of complex diseases, researchers have started to focus on the idea that common disorders are likely influenced by genetic variations that are also common in the population (Reich and Lander, 2001). Instead of looking for individual genes with strong effects on phenotypes, the field has moved toward the study of common, generic variants that individually

have a small effect on the likelihood of having a disease (Bush and Moore, 2012). But what precisely is a variant? At the most fundamental level, two genomes differ in a specific genetic locus if they present an alternative single nucleotide (adenine, thymine, cytosine, or guanine) in that location. Such mutation in one DNA basis is called single-nucleotide polymorphism (SNP) when it appears in at least 1% of the population. One approach for associating SNPs with a disease relies on the fact that a causative variant should be found at a higher frequency in cases than in control subjects. In practice, researchers look for statistical correlations between specific genetic variants and diseases in large samples of people not necessarily related by family ties.

Building over this logic, genome-wide association studies are hypothesis-free methods for identifying associations between genetic regions and traits (Visscher et al., 2017). GWASs compare genetic differences between affected and unaffected individuals using genotyping technologies on large samples. In a typical GWAS project, researchers obtain DNA from two groups of participants, patients with the disease studied and healthy individuals with comparable demographics. Then, selected SNPs on the chromosome are scanned using high-throughput arrays that can genotype up to millions of SNPs for each individual. Variants significantly more likely to appear in the affected patients are biologically crucial for the disease and are likely to be associated with its etiology. Such SNPs might affect gene expressions and functions, mainly when located within a protein-coding gene. It is important to underscore that array-based genome-wide studies do not sequence the DNA base by base, since they only determine the presence or absence of specific SNPs. While microarrays can genotype millions of SNPs, they still constitute a tiny fraction (usually <0.1%) of the genome. Nevertheless, exploiting the fact that co-occurrence of variants at different genetic loci is not random, researchers can use reference genomes (such as the HapMap) to parsimoniously infer the characteristics of the whole genome from the much smaller number of SNPs genotyped (Bush and Moore, 2012).

GWASs have unlocked many significant scientific findings, but have come under scrutiny due to several limitations. First, array-based genome-wide scans can only identify common variants which tag a region likely containing the causal variants of interest. GWAS cannot locate the causal SNPs with certainty, so additional analyses or follow-on studies are usually required to narrow the association region. Second, even when the causal variant is located, the biological mechanism underlying its role in human health requires additional study. Third, the majority of complex diseases are co-determined by a large number of genes, which means that the proportion of variance explained by any individual variant is small. This fact often hinders the therapeutic translation of GWAS findings, since variants with an effect that is too small do not provide actionable drug

targets (Goldstein et al., 2009). Finally, scholars have suggested that GWAS neglects more complex interaction structures between genes, being limited to testing pairwise gene-disease associations (Boyle et al., 2017)

B.3 Hakonarson et al. (2007) and the KIAA0350-Type I Diabetes link

It is estimated that 10% of Americans (around 37.3 million people) have diabetes.⁸ The two main forms of diabetes are type 1 diabetes and type 2 diabetes, with the former accounting for 5-10% of cases. Type 1 diabetes is a chronic condition that often begins during childhood. More in detail, type 1 diabetes is an autoimmune disease that originates from the destruction of β -cells by the immune system (DiMeglio et al., 2018). As a result, the pancreas fails to produce adequate insulin levels, the fundamental hormone needed to allow sugar to enter cells to produce energy and regulate normal glucose levels in the bloodstream. To date, there is no known way to prevent type 1 diabetes, and continuing treatment with insulin is required for patient survival.

Genetics has a sizable role in the emergence of type 1 diabetes: children who have a parent with this condition have a relative risk of 1-9% to present the same condition (DiMeglio et al., 2018). Early candidate gene studies on this disease identified a few genetic determinants, mostly within a set of closely linked genes known as the major histocompatibility complex (MHC). HCM genes code several cell surface proteins that are essential to trigger and target the immune system's response. Malfunctioning in such genes can lead to autoimmune responses, such as type 1 diabetes. Yet, MHC genes explain little more than half of the genetic risk for the disease, indicating that other unknown genetic loci exist. Systematic detection of all remaining genes involved could offer alternative therapeutic pathways and help clarify the etiology of diabetes.

In August 2007, Hakonarson et al. (2007) published an influential genome-wide association study performed on a study population of 563 patients with type 1 diabetes and 1,146 controls.⁹ The analysis was done using a microarray capable of genotyping 550,000 single nucleotide polymorphisms (SNPs). The study identified several SNPs significantly associated with type 1 diabetes. Figure B.1 reports the main results of Hakonarson et al. (2007). Some significant SNPs were located in genes already known to be related to diabetes (e.g., the insulin gene *INS*), but three of them were located in the gene *KIAA0350*.¹⁰ KIAA0350 controls β -cell function and helps prevent

⁸The figure is taken from: <https://www.cdc.gov/diabetes>

⁹The paper also replicated the main analysis on a different sample made of 483 complete family trios, i.e., exploiting differences in DNA between parents and their affected child.

¹⁰The current name for this gene is CLEC16A, but at the time when Hakonarson et al. 2007 published their study it was still known as KIAA0350. For the purposes of this section I will keep referencing it as KIAA0350.

diabetes, but notably it was one of the least studied human genes and its function was unknown at the time of the finding (Soleimanpour et al., 2014).

Figure B.1: Main results from the GWAS analysis of Hakonarson et al. (2007)

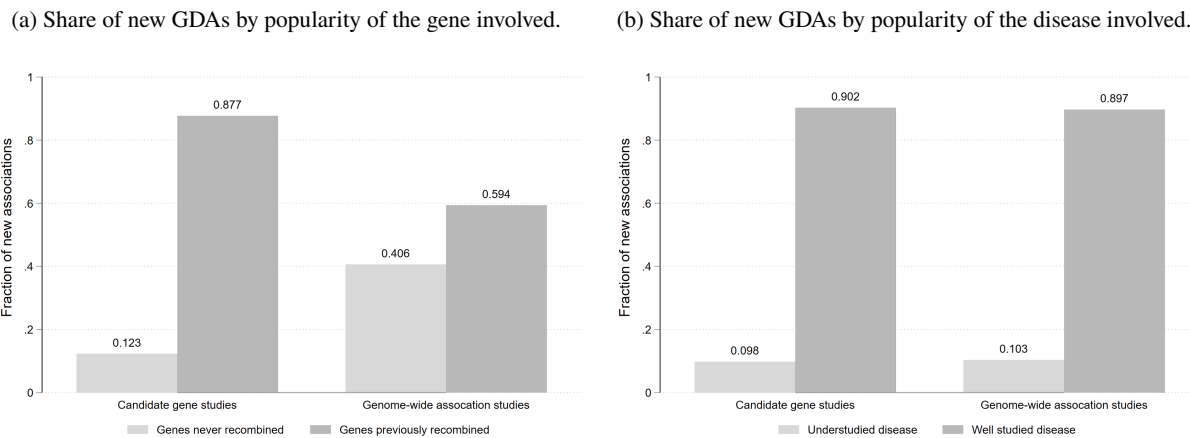
| (a) Results from main analysis | | | | | (b) Text excerpt on KIAA0350 gene | |
|--------------------------------|------------|-------------------|-----------------------|----------|----------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|--|
| Case-control cohort | | | | | | |
| Chr. | SNP | OR (95% CI) | P-value | Locus | | |
| 1 | rs2476601 | 1.80 (1.44, 2.24) | 1.32×10^{-7} | PTPN22 | <p><u>This locus resides in a 233-kb block of LD that contains only KIAA0350 and no other genes, making this gene a prime candidate for harbouring the causative variant. KIAA0350 encodes a protein of unknown function and its genomic location is next to the suppressor of cytokine signalling 1 (SOCS1) gene. The almost exclusive expression specificity of KIAA0350 in immune cells (http://symatlas.gnf.org/SymAtlas), including dendritic cells, B lymphocytes and natural killer (NK) cells, all of which are pivotal in the pathogenesis of T1D^{27,28}, indicates that the variant probably contributes to the disease by modulating immunity.</u></p> | |
| 11 | rs1004446 | 0.62 (0.53, 0.73) | 4.38×10^{-9} | INS | | |
| 16 | rs2903692 | 0.65 (0.56, 0.76) | 4.77×10^{-8} | KIAA0350 | | |
| 11 | rs6356 | 1.52 (1.31, 1.76) | 1.78×10^{-7} | INS | | |
| 16 | rs725613 | 0.67 (0.58, 0.78) | 3.24×10^{-7} | KIAA0350 | | |
| 7 | rs10255021 | 0.58 (0.44, 0.77) | 1.16×10^{-4} | COL1A2 | | |
| 11 | rs10770141 | 0.65 (0.56, 0.76) | 7.20×10^{-8} | INS | | |
| 1 | rs672797 | 1.54 (1.29, 1.85) | 2.67×10^{-6} | LPHN2 | | |
| 16 | rs17673553 | 0.66 (0.55, 0.78) | 1.30×10^{-6} | KIAA0350 | | |
| 11 | rs7111341 | 0.63 (0.53, 0.76) | 3.77×10^{-7} | INS | | |
| 11 | rs10743152 | 0.67 (0.57, 0.78) | 4.73×10^{-7} | INS | | |

Note: Panel (a) shows an excerpt from Table 1 of Hakonarson et al. (2007). The genetic location of the single nucleotide polymorphisms significantly associated with type 1 diabetes are shown in the rightmost column. Highlighted are the three SNPs located in the KIAA0350 gene. Panel (b) shows the passage of Hakonarson et al. (2007) describing the inferred role of KIAA0350. T1D stands for type 1 diabetes.

The KIAA0350-type I diabetes association is robust and has been further investigated in several studies (Gingerich et al., 2020). In my data, this association has a high value of the DisGeNET's GDA Score measure of scientific reliability and no contradictory evidence (Piñero et al., 2020). Twenty-two published papers have experimentally validated the role of KIAA0350 in type 1 diabetes, putting this association in the top 95th percentile of scientific impact. Notably, since the study of Hakonarson et al. (2007), genome-wide association studies have discovered more than 60 additional genetic loci associated with the risk of type 1 diabetes (DiMeglio et al., 2018), setting the stage for an improved genetic understanding of the disease.

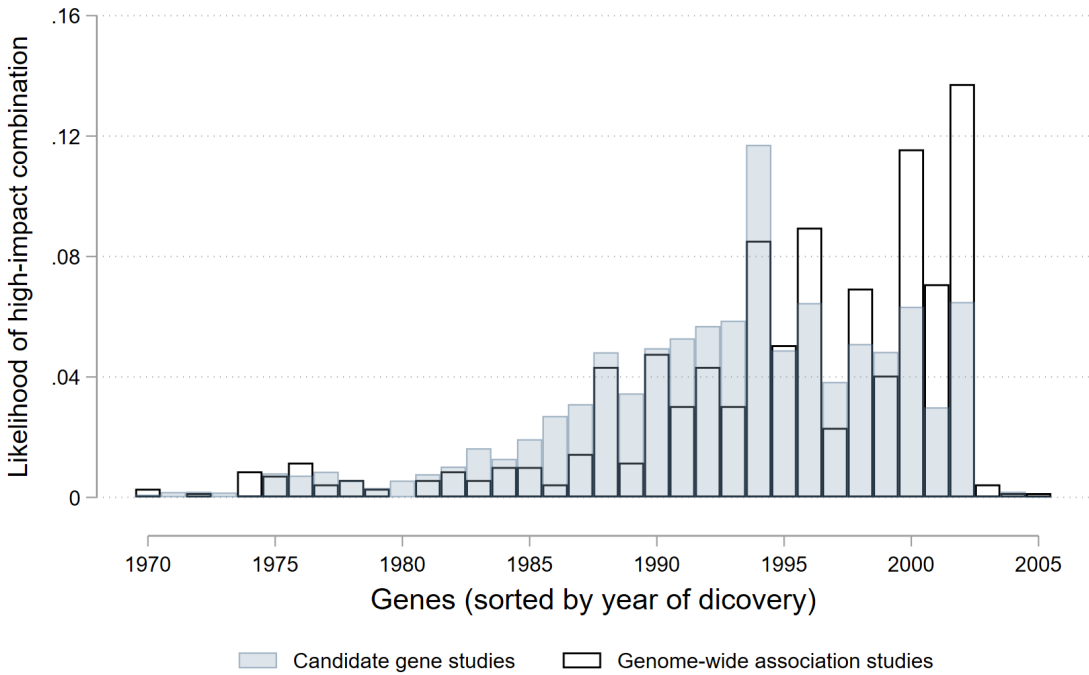
C Additional Figures and Tables

Figure C.1: GWAS induce diversification in gene space but do not change the disease focus.



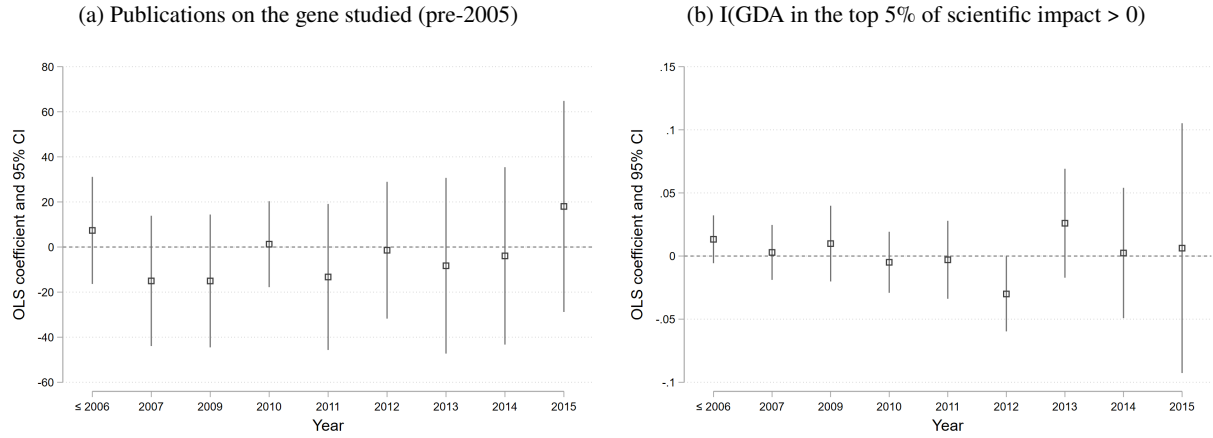
Note: Panel (a) plots the share of new gene-disease associations that involve genes never associated with a disease before vs well studied genes, separately by type of study. Panel (b) plots the share of new gene-disease associations that involve diseases with below median number of gene associations in the DisGeNET data as of 2005 (i.e., less than 2) vs well studied diseases, separately by type of study. Data used in the graph are limited to all new gene-disease associations introduced in the period 2005-2016.

Figure C.2: Gene-disease associations introduced by GWAS are less likely to be high-impact for genes discovered earlier, but more likely for recently discovered genes.



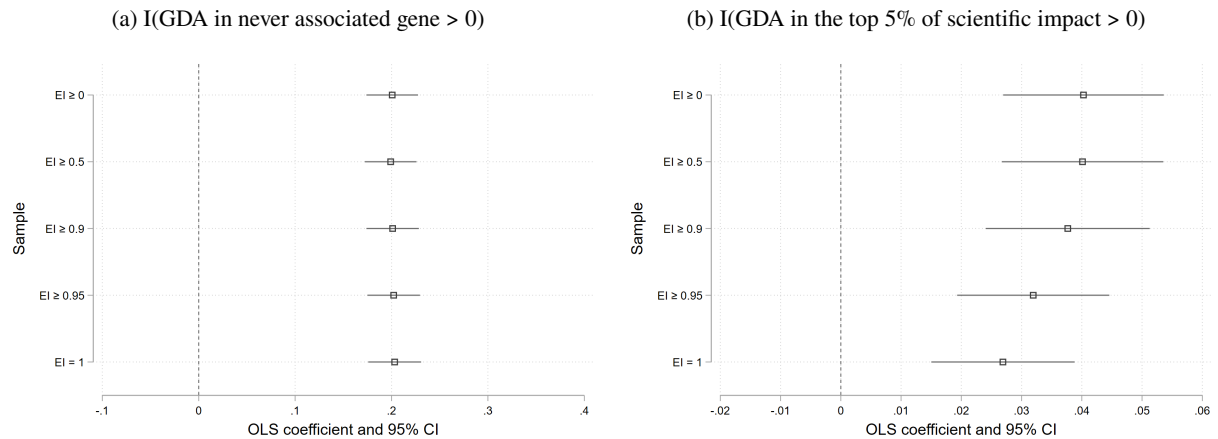
Note: The histogram plots the share of high-impact GDAs for each gene distinguishing by the type of study that introduced them. Data used in the graph are limited to all new gene-disease associations introduced in the period 2005-2016. The 14,112 genes on the X axis are sorted by the year they were first reported in the literature.

Figure C.3: Before their first GWAS, principal investigators that adopt the genome-wide approach are not more likely to target less studied genes or to introduce breakthrough gene-disease associations.



Note: Every coefficient is estimated from a separate regression for a given year, where I compare principal investigators that never publish a GWAS with principal investigators that will eventually publish one but have not done so already (that is, PIs are excluded from the sample after their first GWAS). Panel (a) plots the coefficients and 95% confidence intervals from regressing the popularity of a gene studied over a dummy for whether the principal investigator of the study will eventually publish at least one GWAS. Panel (b) plots the coefficients and 95% confidence intervals of the regression of the likelihood that a GDA is in the top 95th percentile of impact on a dummy that indicates if the principal investigator of the study will eventually publish at least one GWAS.

Figure C.4: Robustness of the main results to the choice of sample.



Note: Panel (a) plots the coefficients and 95% confidence intervals of the regression of the likelihood that gene-disease associations discovered after 2005 involve an understudied gene on a dummy that indicates if the association was introduced by a GWAS. Panel (b) plots the coefficients and 95% confidence intervals of the regression of the likelihood that a GDA is in the top 95th percentile of impact on a dummy that indicates if the association was introduced by a GWAS. In each case the sample is restricted to associations with increasing values of the DisGeNET's *Evidence Index*, which captures the share of contradictory results on the association ($EI = \frac{N_{positive\ pubs}}{N_{total\ pubs}}$). The main analyses of the paper were done on the sample of $EI \geq 0.9$.

Table C.1: Genome-wide association studies are more likely to introduce gene-disease associations involving genes that received less publications or were discovered later than candidate gene studies.

| Dependent Variable: | Pre-2005 pubs on the gene | | Year of discovery of the gene | |
|---------------------------|------------------------------|--------------------------|----------------------------------|-----------------------|
| GWAS | -33.785*** (6.14160) | -30.702*** (10.11909) | 2.159*** (0.20644) | 1.635*** (0.25135) |
| Disease FE | YES | YES | YES | YES |
| Principal Investigator FE | NO | YES | NO | YES |
| Journal prestige FE | YES | YES | YES | YES |
| Year of discovery FE | YES | YES | YES | YES |
| Number of authors FE | YES | YES | YES | YES |
| N | 352,162 | 331,825 | 347,736 | 327,532 |
| Mean of the DV: | 98.886 | 98.886 | 1992.208 | 1992.208 |
| Number of diseases: | 9,740 | 9,362 | 9,663 | 9,281 |
| Number of genes: | 14,072 | 13,910 | 13,770 | 13,615 |

Note: * ** *** denote significance at 10%, 5% and 1% level respectively. Observations at the gene-disease association (GDA) level. Std. err. clustered two-way at the gene and disease level. *Pre-2005 pubs on the gene*: count of the publications received before 2005 by the gene associated to the disease; *Year of discovery of the gene*: year of discovery of the gene associated to the disease; *GWAS*: 0/1=1 for GDAs introduced by a genome-wide association study. See text for details.

Table C.2: Robustness of the main results to alternative definitions of breakthrough discoveries.

| Dependent Variable: | I(GDA in the top 10% of scientific impact>0) | | I(GDA in the top 2.5% of scientific impact>0) | | I(GDA in the top 1% of scientific impact>0) | |
|---------------------------|----------------------------------------------|-----------------------|-----------------------------------------------|-----------------------|---------------------------------------------|-----------------------|
| GWAS | 0.051*** (0.00940) | 0.053*** (0.01112) | 0.032*** (0.00557) | 0.033*** (0.00762) | 0.018*** (0.00389) | 0.017*** (0.00531) |
| Disease FE | YES | YES | YES | YES | YES | YES |
| Gene FE | YES | YES | YES | YES | YES | YES |
| Principal Investigator FE | NO | YES | NO | YES | NO | YES |
| Journal prestige FE | YES | YES | YES | YES | YES | YES |
| Year of discovery FE | YES | YES | YES | YES | YES | YES |
| Number of authors FE | YES | YES | YES | YES | YES | YES |
| N | 350,693 | 330,250 | 350,693 | 330,250 | 350,693 | 330,250 |
| Mean of the DV: | 0.1 | 0.1 | 0.025 | 0.025 | 0.01 | 0.01 |
| Number of diseases: | 9,726 | 9,347 | 9,726 | 9,347 | 9,726 | 9,347 |
| Number of genes: | 12,617 | 12,463 | 12,617 | 12,463 | 12,617 | 12,463 |

Note: *, **,*** denote significance at 10%, 5% and 1% level respectively. Observations at the gene-disease association (GDA) level. Std. err. clustered two-way at the gene and disease level. *I(GDA in the top 10% of scientific impact>0):0/1=1* if the gene-disease association involves a gene in the top 90th percentile of discoveries with the most follow-on work (by year of discovery); *I(GDA in the top 2.5% of scientific impact>0):0/1=1* if the gene-disease association involves a gene in the top 97.5th percentile of discoveries with the most follow-on work (by year of discovery); *I(GDA in the top 1% of scientific impact>0):0/1=1* if the gene-disease association involves a gene in the top 99th percentile of discoveries with the most follow-on work (by year of discovery); *GWAS:0/1=1* for GDAs introduced by a genome-wide association study.

Table C.3: Genome-wide association studies introduce gene-disease associations that receive more follow-on scientific publications than candidate gene studies.

| Dependent Variable: | Count of follow-on publications | | |
|---------------------------|---------------------------------|-----------------------|-----------------------|
| GWAS | 0.545*** (0.19668) | 0.969*** (0.24385) | 1.211*** (0.34976) |
| Disease FE | YES | YES | YES |
| Gene FE | NO | YES | YES |
| Principal Investigator FE | NO | NO | YES |
| Journal prestige FE | YES | YES | YES |
| Year of discovery FE | YES | YES | YES |
| Number of authors FE | YES | YES | YES |
| N | 352,162 | 350,693 | 330,250 |
| Mean of the DV: | 0.56 | 0.56 | 0.56 |
| Number of diseases: | 9,740 | 9,726 | 9,347 |
| Number of genes: | 14,072 | 12,617 | 12,463 |

Note: *, **,*** denote significance at 10%, 5% and 1% level respectively. Observations at the gene-disease association (GDA) level. Std. err. clustered two-way at the gene and disease level. *Count of follow-on publications*: number of scientific articles studying the gene-disease association after it has been introduced the first time by either a GWAS or a candidate gene study; *GWAS:0/1=1* for GDAs introduced by a genome-wide association study.

Table C.4: Gene-disease association studies are more likely to involve the second member of gene families and to find breakthrough discoveries involving them.

| Dependent Variable: | I(GDA with the second member of a gene family>0) | | I(GDA in the top 5% of scientific impact>0) | |
|---------------------------|--------------------------------------------------|-----------------------|---------------------------------------------|-----------------------|
| GWAS | 0.083*** (0.01948) | 0.065*** (0.03060) | 0.036*** (0.01180) | 0.070*** (0.02250) |
| Disease FE | NO | YES | YES | YES |
| Gene family FE | YES | YES | YES | YES |
| Principal Investigator FE | NO | YES | NO | YES |
| Journal prestige FE | YES | YES | YES | YES |
| Year of discovery FE | YES | YES | YES | YES |
| Number of authors FE | YES | YES | YES | YES |
| N | 87,360 | 75,254 | 87,360 | 75,254 |
| Mean of the DV: | 0.39 | 0.39 | 0.05 | 0.05 |
| Number of diseases: | 5,152 | 4,639 | 5,152 | 4,639 |
| Number of genes: | 3,009 | 2,918 | 3,009 | 2,918 |

Note: *, **, *** denote significance at 10%, 5% and 1% level respectively. Observations at the gene-disease association (GDA) level. Std. err. clustered two-way at the gene and disease level. $I(GDA \text{ with the second member of a gene family} > 0): 0/1 = 1$ if the gene-disease association involves the second member of a gene family; $I(GDA \text{ in the top 5\% of scientific impact} > 0): 0/1 = 1$ if the gene-disease association involves a gene in the top 95th percentile of discoveries with the most follow-on work (by year of discovery); $GWAS: 0/1 = 1$ for GDAs introduced by a genome-wide association study. The sample is limited to genes that are members of a gene family.

Table C.5: Diseases targeted first by GWAS are not more likely to be associated to understudied genes.

| Dependent Variable: | I(GDA for never associated gene>0) | | |
|---------------------------|------------------------------------|-----------------------|-----------------------|
| Subsample: | Early GWAS | Mid-period GWAS | Late GWAS |
| GWAS | 0.133*** (0.02652) | 0.108*** (0.02951) | 0.137*** (0.04230) |
| Disease FE | YES | YES | YES |
| Principal Investigator FE | YES | YES | YES |
| Journal prestige FE | YES | YES | YES |
| Year of discovery FE | YES | YES | YES |
| Number of authors FE | YES | YES | YES |
| N | 47,167 | 62,712 | 42,048 |
| Number of diseases: | 111 | 333 | 380 |
| Number of genes: | 9,586 | 9,914 | 8,830 |

Note: *, **, *** denote significance at 10%, 5% and 1% level respectively. Observations at the gene-disease association (GDA) level. Std. err. clustered two-way at the gene and disease level. $I(GDA \text{ for never associated gene} > 0): 0/1 = 1$ if the gene-disease association involves a gene never associated with a disease before 2005; $GWAS: 0/1 = 1$ for GDAs introduced by a genome-wide association study. Column 1 includes only diseases that received their first GWAS before 2009. Column 2 includes only diseases that received their first GWAS between 2009 and 2011. Column 3 includes only diseases that received their first GWAS after 2011.

Table C.6: Robustness of the main results to the use of DisGeNET's GDA Score to measure scientific value of new gene-disease combinations.

| Dependent Variable: | DisGeNET <i>GDA Score</i> for gene-disease associations | | | | |
|---------------------------|---------------------------------------------------------|-----------------------|-----------------------|-----------------------|-----------------------|
| Subsample: | All genes | | | Genes in mouse | Genes not in mouse |
| GWAS | 0.031*** (0.00424) | 0.037*** (0.00410) | 0.038*** (0.00426) | 0.037*** (0.00417) | 0.049*** (0.00920) |
| Disease FE | YES | YES | YES | YES | YES |
| Gene FE | NO | YES | YES | YES | YES |
| Principal Investigator FE | YES | YES | YES | NO | NO |
| Journal prestige FE | YES | YES | YES | YES | YES |
| Year of discovery FE | YES | YES | YES | YES | YES |
| Number of authors FE | YES | YES | YES | YES | YES |
| N | 352,162 | 350,693 | 330,250 | 326,506 | 15,848 |
| Mean if the DV: | 0.06 | 0.06 | 0.06 | 0.05 | 0.06 |
| Number of diseases: | 9,740 | 9,726 | 9,347 | 9,480 | 1,952 |
| Number of genes: | 14,072 | 12,617 | 12,463 | 11,502 | 743 |

Note: *, **, *** denote significance at 10%, 5% and 1% level respectively. Observations at the gene-disease association (GDA) level. Std. err. clustered two-way at the gene and disease level. *GDA Score*= synthetic measure of scientific reliability of the gene-disease association provided by DisGeNET (Piñero et al., 2020); *GWAS:0/1=1* for GDAs introduced by a genome-wide association study.

Table C.7: Genome-wide association studies that introduce new gene-disease associations are more likely to be cited by clinical trial articles than comparable candidate gene studies.

| Dependent Variable: | Cited by a clinical article | | |
|---------------------------|-----------------------------|-----------------------|-----------------------|
| GWAS paper | 0.188*** (0.01994) | 0.193*** (0.01771) | 0.086*** (0.01791) |
| Disease FE | YES | YES | YES |
| Gene FE | NO | YES | YES |
| Principal Investigator FE | NO | NO | YES |
| Journal prestige FE | YES | YES | YES |
| Year of discovery FE | YES | YES | YES |
| Number of authors FE | YES | YES | YES |
| N | 352,162 | 350,693 | 330,250 |
| Mean of the DV: | 0.294 | 0.294 | 0.294 |
| Number of diseases: | 9,740 | 9,726 | 9,347 |
| Number of genes: | 14,072 | 12,617 | 12,463 |

Note: *, **, *** denote significance at 10%, 5% and 1% level respectively. Observations at the gene-disease association (GDA) level. Std. err. clustered two-way at the gene and disease level. *Cited by a clinical article:0/1=1* if the paper that introduced the gene-disease association is later cited by one article describing the outcomes of a clinical trial; *GWAS paper:0/1=1* for papers reporting a genome-wide association study. See text for details.