Does University Licensing Facilitate or Restrict the Flow of Knowledge and Research Inputs Among Scientists?

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Abstract
As university involvement in technology transfer and entrepreneurship has increased, concerns over the patenting and licensing of scientific discoveries have grown. This paper examines the effect of licensing on the citation of academic publications associated with patents covering university scientific research. We analyze data on invention disclosures, patents, and licenses from the University of California, a leading US academic patenter and licensor, between 1997 and 2007. We also develop a novel “inventor-based” maximum-likelihood matching technique to automate and generalize Murray’s (2002) patent-paper “pairs” methodology. We use this methodology to identify the scientific publications associated with University of California patents and licenses.

We find that, in general, licenses are associated with an increase in journal citations to related scientific publications. The timing of this effect supports earlier research that suggests that academic licenses may act as positive signals of research potential in the licensed technological area (Drivas et al. 2014). In contrast, we find the opposite effect of licensing on citations to related scientific publications when the underlying discovery is a research input (which we identify through the use of material transfer agreements (MTAs)). In these cases, the related scientific publications experience a significant decline in citations following the execution of the license.

Our results suggest that, in general, licensing of academic patents does not limit scientific communication linked to patented academic research. Our findings on the effects of licenses on research inputs, however, raise the possibility that licensing may restrict the flow of inputs to further scientific research among researchers.
1 Introduction

Growth in patenting of academic research advances in U.S. and other universities within the OECD has triggered considerable debate since at least 1980, the year of passage of the Bayh–Dole Act in the United States. Supporters argue that patenting and licensing of university inventions can speed the transfer of scientific discoveries to the private sector, promoting the commercialization of such advances. On the other hand, detractors predict a collision between the norms of science and the norms of commerce, fearing that the exclusionary effects of patents will limit research and slow the progress of science. Despite the importance of this question, there has been relatively little empirical work on the extent to which the patenting of academic research results affects scientific research progress. The modest volume of such research (e.g., Murray and Stern (2007)) reflects the challenge of obtaining data that can be used to empirically examine such effects.

This paper develops a new approach to matching scientific publications and patents, which is used to examine the effect of licensing on the journal citations to related publications. We interpret increased citations to these scientific publications after a license issues as evidence that licensing is correlated with a positive effect on the prominence and use by other scientists of the knowledge embodied in the paper, while decreases in such citations may indicate the presence of greater restrictions to such use. Drawing on related work by Walsh et al. (2007), we also examine the effect of licenses on related scientific publications covering the inputs to the experiments of other researchers, a class often referred to as “research tools.”

The next section of this paper discusses the use and potential effects on scientific research of formal intellectual property rights covering academic discoveries. We describe our data and its relevance to this question in Section 3. We then turn to explaining our methodology for constructing publication–patent matches, as well as how we construct a plausible counterfactual for our treatment observations in our methodology (Section 4). We present the results of our analysis (Section 5). Finally, we discuss the implications of our results and conclude with a summary of the contributions and limitations of this paper in Section 6.

2 Research and Intellectual Property in Academia

Universities have long been important performers of research, particularly basic research, in the United States and other industrial economies. The share of U.S. basic research performed by universities has risen in recent years according to the National Science Foundation’s Science and Engineering Indicators (National Science Foundation 2012; Table 4-4).

*** Figure 1 Here ***
At least since the 1970s, this expanded role of U.S. universities in research performance has been paralleled by growth in patenting and licensing of university discoveries, particularly in biotechnology. Figure 2 shows the growth in academia’s share of U.S. patents since 1969.¹

*** Figure 2 Here ***

The roles of universities as sources of basic knowledge and as sources of potentially valuable ingredients for commercial innovation raises the possibility of conflict between these roles, with detrimental consequences for the advance of scientific research. Do patents and licenses restrict access to such knowledge? Or is the existence of a patent and/or license for that patent a signal of the quality of scientific work that leads to greater exploration of the area?

Our examination of the effects of patenting and licensing on scientific communication focuses on “patented publications.” These are discoveries that are published in scientific journals and become the subject of successful patent applications. A patented publication, therefore, is a scientific publication whose discovery is also covered by a patent. Although our patented publications are similar to the patent–paper pairs of Murray (2002) and Murray and Stern (2007), we allow multiple papers to match each patent and thus our observations are not necessarily “pairs.”²

2.1 The Effects of Intellectual Property Protection and Licensing on Scientific Research and Communication

An array of factors, including the Bayh–Dole Act of 1980, other changes in U.S. intellectual property laws and policies, and expanded federal support for academic biomedical research, has increased the patenting of academic research by U.S. universities. The growth in such patenting has been the subject of a large literature and considerable debate over its effects on the scientific research enterprise (e.g., Mowery et al. (2004)).

Heller and Eisenberg (1998) argue that expanded patenting of academic research results may result in fragmented and overlapping property rights covering upstream biomedical research, limiting the ability of scientists to access patented and licensed research results for follow–on research. Other scholars raising concerns over the expanded assertion of property rights in science include Nelson (2003) and David (2003). Empirical research seeking to assess the effects of patenting on scientific communication has examined the effects of patenting on biomedical researchers’ willingness to share information on their work (Blumenthal, et al., 1997; Campbell, et al., 2002). More recent research has analyzed the effects of patenting biomedical

¹USPTO (2012).
²We discuss the construction of our sample of “patented publications” in Section 4.
discoveries that are also disclosed in scientific papers. Some of this work finds that the issuance of a patent results in modest but significant declines in citations to the research papers related to the patent (Murray and Stern 2007, Sampat 2005). Other research, however, argues that biomedical researchers rarely if ever search to determine whether a prospective research project or experiment will infringe on patents (Walsh, et al., 2005; Lei, et al., 2009), raising a question about the mechanism for any observed citation decreases.

Empirical research on the effects of academia’s use of intellectual property has focused mainly on patenting of academic research results. The effects of university licensing on scientific research has received much less attention from scholars. Unlike patents, licenses are not published or otherwise subject to mandatory disclosure. In many cases the identity of licensees is treated by university technology transfer offices as confidential (Ziedonis 2007).

Why and how might licenses affect the behavior of academic researchers in formulating their research agenda? Sampat and Ziedonis (2005) examined patent citations to Columbia University and University of California patents that were licensed. They found that citations were associated with an increase in the likelihood that a patent would be licensed. Moreover, most citations occurred after the patent was licensed. These scholars interpret this pattern as indicating market interest in the technological area surrounding the licensed patents. More recently, Drivas et al. (2014) found that citations by non-licensees to patents exclusively licensed (either by geographic area or field of use) by the University of California increased after the licenses were executed. Similar to Sampat and Ziedonis, Drivas et al. regard the increase in non-licensee patent citations as a reaction to the potential commercial value signaled by the negotiation of the license.

It is possible that a similar signaling effect associated with the execution of a license could increase citations to patented publications linked to the license. In such a case, the issue of a license “demonstrates” that a particular area of research has potential scientific or commercial value, leading other investigators to pursue work in closely related fields. It is also possible (see Larsen (2011)) that contemporary academic researchers may choose research areas partly based on their potential private profitability, and therefore might respond positively to a “signal” that a given area of research has attracted the attention of industrial licensees. Regardless of whether a license signal operates through perceptions among researchers of scientific or commercial potential, this argument predicts an increase in citations to patented publications following the negotiation of the license.

Equally plausible arguments, however, suggest a chilling effect of licensing on scientific communication. Reactions by university technology licensing offices and/or their licensees to any evidence of patent infringement (even for research purposes, inasmuch as the research exemption from such infringement suits remains unclear) may be swifter and stronger in the case of patents that are licensed. And licenses may include provisions for reach-through royalties and limitations on the disposition of intellectual property on follow-on
research. Moreover, the negotiation of a license may take considerable time, delaying access to the materials or tools embodied in the disclosure.

We are thus agnostic on the likely direction of any effect of licenses on scientific communication associated with publications linked to licensed academic patents. Indeed, both effects may be present for papers in various fields of research, and we hope that our findings shed light on the magnitude of any offsetting effects.

2.2 Research Inputs and Material Transfer Agreements

Patents and licenses increase the “excludability” of intellectual property for other researchers, exposing them to potential legal liability in the event that they utilize or exploit the intellectual property protected by patents and issued to others for their own research or commercial use. A very different form of “excludability,” highlighted by Walsh et al. (2007), concerns the denial by one researcher of physical access to materials (or other research results) that are inputs to the experiments of another researcher.\(^3\) The survey results of Walsh et al. (2007) indicate that such denials can impose significant costs and delays on the scientific work of other researchers, costs and delays that according to these authors, exceed those associated with patents.

Research inputs have been widely (and imprecisely) identified as “research tools.” For example, the NIH Working Group on Research Tools (1998) defines them as “the full range of resources that scientists use in the laboratory… the term may thus include cell lines, monoclonal antibodies, reagents, animal models, growth factors, combinatorial chemistry libraries, drugs and drug targets, clones and cloning tools (such as PCR), methods, laboratory equipment and machines, databases and computer software.”\(^4\)

As we noted above, Walsh et al. (2007) argue that denials of access to such research tools are more likely when the erstwhile supplier of them is engaged in “commercial activity,” such as licensing of the invention disclosure associated with them. Interviews we conducted for this study supports this view, with one scientist (who was involved in a start-up firm) saying “if another company asked to use our [materials] for [same purpose as our company uses them] we would say ‘no.’”\(^5\)

Even when access is granted, however, researchers gaining access to research tools that are associated with licensed disclosures may encounter difficulties, as highlighted by the celebrated case of the Oncomouse (Murray 2010):

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\(^3\)“More problematic [than patents in limiting researcher access] is access to materials and/or data possessed by other researchers, such as cell lines, reagents, genetically modified animals, unpublished information, etc. Restrictions on access, however, do not appear to turn on whether the material is itself patented. Rather, such restrictions are more closely associated with scientific competition, the cost of providing materials, a history of commercial activity on the part of the prospective supplier, and whether the material in question is itself a drug.” (Walsh et al. 2007; p. 1185)

\(^4\)The National Research Council panel has a similarly broad definition that includes materials that “…may be critical inputs for the success of a research project.” (National Research Council 2010; p. 7).

\(^5\)Even though this scientist was unwilling to share materials in this instance, in other instances he/she had shared materials. This is important for this study, because some successful transfers of materials can be observed.
“In 1984, scientists at Harvard University carefully engineered a new mouse to have a predisposition to cancer, the Oncomouse... The Harvard researchers... patented their creation and subsequently licensed this patent to DuPont... [Dupont] set a high price per mouse... placed restrictions on breeding programs... demanded publication oversight... [and] insisted upon a share of any commercial breakthroughs made using the Oncomouse.”

These and similar restrictions may have an adverse effect on the use by other researchers of research tools such as the Oncomouse. Moreover, as the NIH Working Group on Research Tools (1998) noted in its report, licensees may have an incentive to restrict access to these materials:

“If the sponsor or licensee plans to develop the research tool as a commercial product for sale to researchers, it may be unwilling to permit the university to undercut its position in this particular market by giving the tool away to potential paying customers.”

The surveys of scientists by Walsh et al. (2007) and Lei et al. (2009) found requests by researchers for research tools from industrial researchers, a group more likely to be engaged in commercial applications of research, are rejected approximately twice as often as requests to other academics.

Based on these arguments, we anticipate that access to research tools (inputs to other scientific experiments) that are associated with licensed intellectual property (a sign of the commercial exploitation, prospective or otherwise, of the intellectual property) may well be restricted, even when terms for its exchange among researchers are successfully negotiated through a Material Transfer Agreement (MTA). Licensing of IP related to research tools thus may have negative consequences for follow-on scientific research and therefore may have a negative effect on citation rates for publications related to such IP.

As the discussion of “research tools” thus far indicates, developing a definition that facilitates their identification and empirical analysis is challenging. The definitions of research tools employed by the NIH Working Group and Walsh et al. (2007) are very broad and do not lend themselves to empirical operationalization. Instead of attempting to develop and defend a definition of research tools that relies on the characteristics of the relevant invention disclosure or patent, we identify research tools based on the association of a Material Transfer Agreement (MTA) with a given patented invention disclosure.

Material Transfer Agreements (MTAs) are agreements that govern the transfer and exchange of materials, usually biological, used in research. Although the informal exchange by researchers of biological materials for use in fundamental research has a long and occasionally controversial history in the biomedical sciences, these materials exchanges historically were governed by little more than a letter from the source accompanying the materials, requesting acknowledgement and in some cases asking that the materials not be passed on to third parties (McCain 1991). The more elaborate MTAs used in contemporary materials exchanges appear to be a byproduct of the post–1980 surge in academic patenting (Streitz and Bennett 2003). One of the few
analyses of the role of MTAs in the scientific research enterprise is Stern’s discussion of biological resource centers (2004).

Biological resource centers (BRCs) are nonprofit materials depositories that play a key role in maintaining the reliability and provenance of cell lines used by industrial and academic researchers—as Stern notes, contamination of widely used cell lines has caused major research fiascoes in the past several decades. Stern argues that the use of MTAs by BRCs has aided the exchange of materials, and recommends that MTAs be a standard complement to patents covering biological discoveries: “Putting MTAs in place at the time of patent approval lowers the cost of mutually beneficial transactions between the developers of materials and follow–on researchers and widens the availability of patented biomaterials” (2004; pp. 96–97). Similarly, Walsh et al. (2003) argue that the formalization of materials exchanges through MTAs may simplify these transactions and facilitate researcher access.

To confirm that MTAs are a good indicator that an invention disclosure is associated with a research tool, we examine U. C. Berkeley data on incoming MTAs—that is, research materials requested from other researchers by U.C. Berkeley researchers. These data describe the requested materials and their intended use by the recipient U. C. Berkeley researcher. We analyzed a random sample of 50 of these MTAs, and found that 44% were related to DNA/RNA/Plasmids, 32% concerned cell lines or other biological/chemical agents, 16% were animal models, 6% were data transfers, and 2% were concerned with “other non–research inputs.” Overall, therefore, 98% of these MTAs involved materials that fit within the NIH Research Tools Working Group definition of research tools. We also analyzed the intended use for the materials requested through the MTA by U. C. Berkeley researchers, and found that 94% of the MTAs indicated that the requested material was to be used as an input to further research and a further 4% implicitly indicated that such uses were intended.

Based on our analysis of U. C. Berkeley MTAs therefore, we believe that our treatment of the presence of an MTA as an empirical indicator that a given invention disclosure is indeed a research tool and/or input to follow–on scientific research is defensible. It is important to note two caveats associated with our empirical use of MTAs as indicators of research tools, however. As Walsh et al. (2007) and Mowery and Ziedonis (2006) point out in their discussions of MTAs, a majority of the materials transfers among academic scientists do not rely on formal MTAs that are disclosed to academic Technology Transfer Offices (TTOs). Although we believe that the presence of an MTA is a reasonable indicator that a given disclosure has applications as a research tool, our data in fact contain many other disclosures (including disclosures that are patented and licensed) that may well be research tools but (lacking an MTA) cannot be identified as such. It is likely that our empirical approach thus understates the effects on citations of licensing of patented disclosures that are research tools. In addition, our data enable us to only identify the “effects of MTAs” that are negotiated
and agreed to by all parties to the materials transfer. In other words, and in contrast to Walsh et al. (2007),
we do not identify the effects on scientific research of the denial by researchers of other researchers’ requests
for research materials. Nevertheless, our results seem to indicate that the types of restrictions associated
with MTAs linked to licensed disclosures that were examined by Murray (2010) in her discussion of the
Oncomouse may indeed have effects on scientific communication, as measured by citations to publications.

3 Data

We draw on two principal sources of data for our empirical analysis. The first, the “IP data,” is an extract
from the technology disclosure database maintained by the Technology Transfer Office within the University
of California Office of the President (UCOP). UCOP monitors and in some cases manages invention disclo-
sures, patent applications, and licensing transactions for all campuses of the University of California (nine
campuses, including five medical schools, during the period of our study).

These data list all 11,341 inventions reported by University of California faculty from 1997 to 2007. These
disclosures led to 2,035 issued U. S. patents, 1,890 licenses to these patents, and 3,853 MTAs by the end
of 2009. Note that only a small subset of technology disclosures is patented, and universities’ patenting
propensity varies among fields of academic research—since the 1980s, patenting and licensing activity at
U. C. has been dominated by biomedical research (Mowery, et al., 2001). The distribution of MTAs also
is highly skewed, with few disclosures generating the majority of MTAs and many disclosures associated
with no MTAs. This echoes the finding in Mowery and Ziedonis (2006) that MTAs are disproportionately
concentrated in biomedical fields of research, as are licenses.

The second source, “publications data,” comes from Web of Science, an internet–based service that tracks
the bibliographic information and the citations to and from articles published in 10,000 of the highest-impact
journals across 256 disciplines.6

From Web of Science we gathered the title, author names, journal, publication date, and citation infor-
mation for each scientific paper. The information on “forward citations,” citations from later published
papers to that publication, was extracted through the end of 2009. Web of Science also provides a number
of well–accepted measures of journal quality. The most prominent of these is the “impact factor,” which
measures the average number of times an article in that journal is cited in its first two years, and which we
include in our analysis.

4 Methodology

4.1 Sample

Our study focuses on finding differences in the citations associated with patented publications, depending on whether or not they are licensed. One advantage of restricting our comparison to patented publications is that they are more likely to be similar in quality and other characteristics. In particular, because all of the underlying disclosures are patented, differences in “commercializability” are considerably lower than they would be in a general sample. Nevertheless, there may well exist other unobserved differences within our sample between the patented publications that are licensed and those that are not. Below we describe how we construct a control sample to address these issues.

Our sample is also restricted to patented publications with at least three authors, which is a by–product of the method that we use to match publications and patents (discussed in detail below). This restriction imposes two additional conditions on our sample. First, a patent must have at least three inventors. Second, the associated publication must list at least three of those inventors as authors. Figure 3 depicts the consequence of these restrictions on the sample.

*** Figure 3 Here ***

Column 1 in Figure 3 represents the 2035 patents in the full sample. By excluding patents with fewer than three inventors, we omit 944 patents listing one or two inventors (Column 2) from the full sample, leaving 1091 patents. From these 1091 we exclude an additional 363 three–or–more–inventor patents where fewer than three inventors were listed as authors on any publication, resulting in a remaining sample of 728 patents (Column 3). Of these 728 patents, 406 list three inventors, 201 list four inventors, and 121 list 5 or more inventors. The fourth column of Figure 3 reports the number of journal citations “per patent,” (i.e., the number of journal citations for all publications that are matched, using the three–name overlap restriction, to that patent). Column 5 contains 716 patents with citations and a further 12 with no citations (labeled as “partially in the sample”). These latter 12 are kept in our sample to avoid excluding them based on the outcome variable (moreover, they are at risk of being cited). The resulting sample consists of 728 patents.

At the other extreme, a few publications receive large numbers of citations, creating the risk that these outliers would unduly influence our empirical results. To ensure that the results are not driven by outliers, we trim the 2.5% highest and lowest treated observations and then examine the remaining distribution of residuals.
4.2 Matching Invention Disclosures, Patents, and Scientific Publications

4.2.1 Constructing Patent–Paper “Pairs”

Our empirical analysis adopts a methodology that is broadly similar to that of Murray and Stern (2007). We measure “scientific communication” as the number of times a published scientific paper is cited by subsequent articles published in scientific journals (a “count”). Our sample includes only scientific publications that are patented—that is, where the discovery has also been disclosed by the researcher to the university, and the university has subsequently patented it. We also observe the licenses and MTAs, if any, associated with these patented invention disclosures. To construct this sample, we rely on the connections between patents, licenses and MTAs provided by UCOP. Connecting patents to scientific publications is less straightforward. Murray and Stern (2007) matched patents to articles published in the journal *Nature Biotechnology* by reading both patents and the academic articles and relying on expert judgment to link them. Our methodology instead employs an “inventor–based matching” technique to link patents to scientific papers.

Inventor–based matching relies on two assumptions. First, inventors listed on a patent are likely to be the authors listed on related publications. Second, the patent application date is likely to occur near the publication date of the academic article. Based on these assumptions we construct a maximum–likelihood estimator for the publication(s) that best matches a particular patent. For each inventor name listed in the patent, we first identify all publications authored within a five year window that includes the year of the patent application and the two years prior to and following that year (\(t-2\) to \(t+2\)). We then determine the instances in which the inventors’ publications overlap. Those publications with the greatest overlap are chosen as matches. For example, if a patent has three inventors, we extract three publication sets (one for each inventor) and retrieve publications common to all three inventors. Figure 4 illustrates this process for the three–inventor case.

*** Figure 4 Here ***

This approach can result in multiple publications as “best” matches, in contrast to Murray and Stern (2007), which matches one publication to each patent. To illustrate, consider patent number 7011723, “Adhesive Microstructure and Method of Forming Same.” The patent pertains to adhesives inspired by the design of gecko feet and credits four inventors. The inventor–based matching approach defined above yields two matches of the underlying invention to publications in scientific journals. The first, entitled “Adhesive Force of a Single Gecko Foot–Hair,” appeared in *Nature* in March 2000. A second article, “Evidence for Van der Waals Adhesion in Gecko Setae” was published in *Proceedings of the National Academy of Sciences* in 2002. In this case, all four patent inventors were listed on each publication.
Inventor–based matching does not restrict us to instances in which all inventors are listed as authors on the publication. For instance, if in the example above a lab technician had also been listed as an inventor on the patent, but was not included on any linked academic publications, the algorithm would choose the publication(s) with the maximum possible overlap. In this instance the publications listing four of the five inventors would be chosen since there would be no five–out–of–five–inventor matches.

4.2.2 Maximum–Likelihood Estimation for Patent–Paper Pairs

As highlighted above, an implication of the inventor–based matching method is that a single patent can be associated with more than one publication. Such a match will occur precisely when multiple publications share the same level of overlap between the inventors, and when no publications have a greater overlap. Based on the “best–available” property of maximum–likelihood estimators, we assume that a publication and a patent are more likely to be a match if they share an author:

\[
p \left( \text{match}_\text{pub}_i \cap \text{patent}_j | \text{authors} \subset \text{authors}_{\text{pub}_i} \cap \text{inventors}_{\text{patent}_j} \right) > p \left( \text{match}_\text{pub}_m \cap \text{patent}_j \right)
\]  

Here \( \text{authors}_{\text{pub}_i} \) and \( \text{inventors}_{\text{patent}_j} \) are the sets of authors for publication \( i \) and the inventors for patent \( j \), respectively. Given these parameters, \( \text{pub}_m \) is a “match” for \( \text{patent}_j \) if

\[
m \in \arg \max_i \prod_{i,k} p \left( \text{match}_\text{pub}_i \cap \text{patent}_j | \text{authors} \subset \text{authors}_{\text{pub}_i} \cap \text{inventors}_{\text{patent}_j} \right)
\]  

As with all maximum–likelihood estimators, a “best” estimate is not necessarily precise (Casella and Berger 2002). In general, accurate matches are less likely when the precision of the estimate is low. For example, our matching algorithm would be unlikely to produce correct matches if, on a four–inventor patent, we identified only publications that listed a single inventor as an author. In this case, the algorithm would theoretically identify as matches all publications by all of the inventors in the relevant \( (t-2, t+2) \) window. To avoid such errors and improve the precision of the estimate, we restrict matches to those with three or more authors. The logic behind this criterion is illustrated in Figure 5, which portrays the number of papers matched to each patent in our dataset linked by at least two inventors and authors and lists the number of names common to both the patent and the published paper.

*** Figure 5 Here ***

Figure 5 shows that 82% of the sample patents is linked by the inventor–based matching algorithm to between one and five papers, while the remaining 18% of our patents matches to six or more publications. The
large number of papers associated with each of the patents in the 18% may reflect common scientist names (e.g., “J. Smith”). Figure 5 also demonstrates that patents matched to more than five publications typically have only two names that are common to both the inventor list and the author list. We therefore exclude all the patents marked “2” in Figure 5, noting that we have previously excluded patents and papers with only one name in common. This leaves us with 728 patents and their related publications. The statistical implication of dropping publications with an overlap of two authors is to restrict the sample to patent–paper pairs with a higher expected probability of a match, thus reducing the likelihood of including false matches (hereafter “noise”) in our sample.

We examine the effect of restricting our sample to higher levels of inventor–overlap by comparing the sample statistics of 3–inventor overlap and 4–inventor overlap samples, as shown in Columns 1 and 2 in Table 1. Column 3 reports difference of means test results for each variable.

*** Table 1 Here ***

The average publication year, publication age when the citations are observed, and the proportion of the papers in the Life Sciences (principally biology, biochemistry, and medicine) are relatively stable across the samples, suggesting little or no introduction of bias along these dimensions.\(^7\) In contrast, two measures of publication quality, the number of citations per year and the average impact factor of the publication’s journal, are higher in the 4–inventor overlap sample in Column 2. Since correct matches are patented publications by definition and incorrect matches are random additions from the general pool of publications, it is not surprising that the removal of incorrect matches yields an increase in average publication quality. This result is consistent with our argument that higher–overlap specifications reduce noise in the sample.\(^8\) It is also consistent with Murray and Stern’s (2007) finding that patented publications receive more citations than unpatented publications.

4.2.3 Validation of the Inventor–Based Matching Technique

To assess the validity of this matching algorithm, we compare its output with the hand–matches compiled by Murray and Stern (2007).\(^9\) For each patent in the Murray and Stern sample, we generated maximum–likelihood estimates of the “best” publication matches using the inventor–based matching technique. Of the 170 patent–publication matches identified by Murray and Stern, our automated method determined an identical “best” publication match for 95% of the sample. In a small number of cases (4%), our algorithm

\(^7\)As Table 1 shows, while small, some of these differences are statistically significant. In the case of Publication Year, the difference is equivalent to a few months.

\(^8\)All else equal, it is plausible that publications associated with patents will tend to be of higher quality than an average unpatented publication. So, if tightening the “overlap” restriction results in a sample with fewer unpatented publications (“noise”), then it should have a higher average citation rate, which is what we observe.

\(^9\)Fiona Murray and Scott Stern graciously provided their data to us for comparison.
identified a “better” match. Only in two cases (1% of their sample) did our approach yield matches otherwise inconsistent with their hand-matching process.

In summary, our inventor-based matching approach, while demonstrating accuracy comparable to the hand-matching approach employed by Murray and Stern (2007), possesses several advantages, (a) it does not impose a simple one-to-one relationship between patents and publications, (b) it is transparent, reproducible, and does not rely on domain expertise, (c) it is applicable to multiple scientific fields, and (d) it is automated, making larger sample sizes feasible.

4.3 Construction of Treatment and Control Groups

Designing an adequate specification to test the effect of licensing on scientific communication is a challenging exercise, because of the complex and varied patterns of citations that publications may receive during our sample period. Citations to some publications may grow throughout the period of our data, whereas for other publications, citations may rise, peak, and then decline. Moreover, the timing and rate of the ascent and descent in citations may vary. It is thus difficult to construct a sufficiently flexible parametric model to adequately accommodate these differences. Moreover, if the specification is not sufficiently flexible, a poorly modeled sub-group could bias our results. Because of the complex and potential non-linear interactions of various control variables that we may employ to construct treatment and matching control groups, we therefore pursue a flexible nonparametric approach. This technique allows us to weaken the assumption of linearity and enables us to account for the many plausible interaction effects that could be present in our analysis.

The non-parametric method we use is “nearest neighbor” matching. It searches the set of non-treatment observations to identify the one that “best” matches each treatment observation. Collectively, these “best matched” non-treatment observations form a control group. Because this search is done based on observable characteristics, the control and treatment observations should be similar along these dimensions by construction.12

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10 The difference here is likely because of the direction of matching. Murray and Stern began with a set of publications and found the most-similar patent, whereas our analysis starts with a patent and finds the most-similar publication. The patent identified by Murray and Stern may have been the best match for that publication, but another publication may be an even better match to that patent. Our non-expert review of the “better” matches suggests that they are well matched.

11 It is reasonable to assume that publications in more highly cited journals accrue more citations, which would argue for including Journal Impact Factor as a control variable. Similarly, the academic discipline (hereafter Journal Subject) could also drive citation patterns, as could how much higher cited a publication is prior to the license (Citations in t−1, Citations in t−2). Each of these the effects could be non-linear, which would suggest the inclusion of higher-order terms. Interaction terms between these variables would also be important, since the effects of Journal Impact Factor could differ by discipline. Each of these effects would affect a publication differently at different points in time, calling for them to be interacted with age fixed effects. Fully interacting all of these effects to capture the entire complexity would result in many of the parameter estimates to be determined by small numbers of observations, or have no observations at all.

12 Restricting ourselves to control observations that match the treatment observations decreases our sample size. The effect that this has on the precision of our estimates is ambiguous, however. Smaller samples will tend to reduce our precision (making the standard errors larger), but there may also be a countervailing effect as the smaller sample will likely be more homogenous,
Since matches are found for the treated observations, the estimate from our analysis should be interpreted as an “average treatment effect on the treated” (ATT), that is, the average effect of a license on citations to patented publications similar to those that do receive licenses (in contrast, for example, to the effect of a license on an average publication or an average patented publication).

We identify the control matches using the “Genetic Matching” technique, which is a generalization of the propensity scoring matching method (Rosenbaum and Rubin 1985) that employs a search algorithm to iteratively improve covariate balance (Diamond and Sekhon 2013, Sekhon 2011). Our procedure employs two types of variables, those where we require an exact match and those where a nearby (“nearest neighbor”) match is sufficient. We force an exact match on the following variables:

- Publication Age: Number of years since the paper was published
- Journal Subject: Academic discipline of the journal (e.g., medicine)
- Patent Granted (yes/no): Whether the related patent has been granted at the time of license
- MTA Issued (yes/no): Whether the paper has an associated MTA at the time of the license

For example, these restrictions imply that in our analysis a (licensed) treatment observation in the life sciences with an issued patent and no MTA would be compared with a control group observation in the life sciences with an issued patent and no MTA, and that the time period covered by the comparison would be identical for the treatment and control observations.

For each treatment observation that matches on these exact characteristics, we then choose its nearest neighbor based on its relative proximity in the following five characteristics:

- Journal Impact Factor
- Publication Year
- Citations in \( t - 2 \): Number of citations two years before the treatment
- Citations in \( t - 1 \): Number of citations one year before the treatment
- Slope of citation curve between \( t - 2 \) and \( t - 1 \).

In each of these dimensions we limit the maximum “distance” between each treatment and control observation to one standard deviation for that variable (the “caliper”), thus excluding any observation that differs and thus increase precision.

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13 Treatment and control variables are in “covariate balance” when they have the same joint distribution of their observed covariates (Diamond and Sekhon 2013). According to Sekhon (2011) in referring to the genetic matching software GenMatch, “GenMatch dominates the other matching methods in terms of MSE [Mean Squared Error] when assumptions required for EPBR [Equal Percent Bias Reduction] hold and, even more so, when they do not.” Previous social science researchers who have also used this generalization of the propensity score matching method include Morgan and Harding (2006), Gilligan and Sergenti (2008), Eggers and Mainmueller (2009), Ladd and Lenz (2009), Gordon (2009), and Hopkins (2010).
by more than that amount on any characteristic. Under this procedure, treatment observations with no equivalent control observation are dropped from the sample.

We further control for any remaining differences in the characteristics between treatment and control observations by applying a “covariate bias adjustment.” We conduct a multivariate linear “regression adjustment” on the post–matching sample of treatment observations plus their matched controls. Collectively, these restrictions result in a control group such that for each characteristic, the observation matches either exactly or within one standard deviation to the corresponding treatment observation characteristic.

### 4.4 Methodology

Once we have generated our treatment and control group sample, we employ a difference–in–differences approach to estimate the size of the treatment effect. More specifically we compare the change in the number of citations to one patented publication following the execution of a license (a treated observation) to the change in number of citations for a comparable publication that lacks a license (a matching control observation). The specification is as follows:

$$Treatment\ Effect = (\text{Citations}_{t+1} - \text{Citations}_{t-1})_{pub/w\ License} - (\text{Citations}_{t+1} - \text{Citations}_{t-1})_{pub/w/o\ License}$$  \hspace{1cm} (3)

Using a differences–in–differences estimator allows us to avoid bias associated with changes that affect the “before” and “after” periods simultaneously. For example, in the matched sample case, if a particular scientist receives on average five additional citations (relative to the sample average) per year, these citations will be included in both \((\text{Citations}_{t-1})\) and \((\text{Citations}_{t+1})\) terms, thus the impact on the estimate will be zero. Our estimator thus accomplishes the same effect as author or publications fixed effects. It is also robust to these types of unobservable differences, as well as the observable differences controlled for using matching.

To summarize, we obtain our covariate balance by using two techniques. First, we use matching to find the nearest neighbor to our treatment observations. This non–parametric technique means that we make fewer assumptions about the parametric form of the effect than we would with a linear, or generalized linear (e.g., negative binomial) formulation. Secondly, we perform a multivariate linear regression (using the same covariates used in matching), to adjust for any remaining differences between the groups. \footnote{Uncoarsened exact matching (CEM) is an alternative matching approach. Employing a caliper rather than CEM, however, allows us to exclude both observations whose observable covariates would make them outliers and those which would qualify them as “inliers,” that is, observations that are within the range of the data, but nevertheless lack a comparable control observation (Sekhon 2011).}

\footnote{For the license effect: \(\Delta\text{Citations} = \beta_0 + \beta_1\text{Age} + \beta_2\text{Journal Subject} + \beta_3\text{Patent Granted} + \beta_4\text{MTA Issued}_{Y/N} + \beta_5\text{Journal Impact Factor} + \beta_6\text{Publication Year} + \beta_7\text{Citations}_{t-2} + \beta_8\text{Citations}_{t-1} + \beta_9\text{Citations Slope} + \epsilon.\) Our coefficient...}
expect such differences to be smaller, the linearity assumption embedded in least-squares is more plausible, although still subject to the same criticisms noted above. Rubin (1979) discusses the value of using these (slightly modified) techniques and concludes that “pair-matching coupled with regression adjustment on the matched pairs is a quite effective general plan for controlling the bias due to matching variables, and this combination is clearly superior to regression adjustment” (p. 318).

5 Results

Before we estimate the effects of licensing on citations to patented publications, we first must verify the covariate balance between the treatment and control groups. An effective matching procedure should yield summary statistics for the treatment and control groups that are similar. Table 2 reports the mean of each group for the control variables and the results of $t$-tests (difference of means) and Kolmogorov–Smirnoff (KS) tests (difference in distributions).

*** Table 2 Here ***

The equal means (and equivalent distributions) across the treatment and control groups for first four variables reported in Table 2, Publication Age, Journal Subject, Patent Issued, and MTA Issued, reflect our requirement of exact matching on these dimensions across the two groups. For the remaining variables, means for the control and treatment groups appear to be similar, but some differences are statistically significant. Most notably, the means and distributions between the two groups are significantly different at a 1% level for Journal Impact Factor and Citation Slope from $t-2$ to $t-1$. The Kolmogorov–Smirnoff test also shows that the distribution of Citations in $t-1$ differs at 1% significance across the two groups. Overall, the presence of significant differences in several of the matching variables suggests that the regression-adjusted analysis improves covariate balance beyond that achieved by relying solely on the caliper matching approach.

5.1 The Effects of Licensing on Citations to Patent-Linked Publications

We now turn to estimating the effects of licensing on scientific communication as measured by citations to patent-linked publications.\textsuperscript{16} We report the results of our findings in two ways. The effect can be seen directly through the citation pattern for the treatment group (dashed line with circles) and the control group (dotted line with squares), as shown in Figure 6.

*** Figure 6 Here ***

\textsuperscript{16}Although exclusive licenses represent 97% of the licensing activity (with 3% of the licenses being non-exclusive), results are reported by types of license. An analysis of exclusive licenses only (not reported) produces similar results.
As depicted in Figure 6, patented publications receive more citations starting two years after a license is executed than do publications that are linked to unlicensed patents. This pattern of increased citations two years after the license is consistent with a more gradual expansion of awareness within the research community of the license that increases citations to the relevant publication only after a lag, reflecting the lack of any public announcement of the license. In this interpretation, scientists gradually adjust their research agendas to increase their attention to the area of research at the center of the license, in response to the positive signal of the commercial or scientific quality embodied in the license.

Figure 7 reports results from the regression adjustment analysis, confirming the post–licensing citation pattern presented in Figure 6 and testing for the statistical significance of the differences in citations between the two groups. The effect of licensing on the number of citations is near zero and not statistically significant in the first year after the execution of the license. The average difference in citations between the treatment and control groups is 2.4 in the second year and 2.1 citation in year \( t+3 \), both of which are significant at the 1% level. This pattern is very similar to the citation trends depicted in Figure 6 due to the covariate balance prior to the regression adjustment. The magnitude of the differences between the two groups in years \( t+2 \) and \( t+3 \) implies that the average publication receives an increase of approximately 25% in citations in these two years compared to the control group.

*** Figure 7 Here ***

Inspection of the residuals after our regressions (not reported) shows that the mean and median are aligned and that the residuals are distributed approximately normally, supporting the validity of our model.

5.2 The Effects of Licensing on Citations to Patent–Linked Publications Associated with Research Tools

We now turn our attention to the effects of licensing on patented publications that we believe are more likely to be associated with research tools. In their test of the effects of patenting on citations to “research tool”–related publications, Murray and Stern (2007) defined such publications as those linked to any patents within their sample in the 3-digit patent classes 435 (Chemistry: Molecular Biology and Microbiology) and 800 (Multicellular Living Organisms and Unmodified Parts Thereof and Related Processes). A patent class–based definition, however, does not account for the patent’s use. Instead of employing patent classes, as we noted above, we believe that patented publications for which MTAs are issued are likely to fall within a rough definition of “research tool.”

Our analysis of the effects of licensing on this class of patented publications restricts the sample to only
those with materials transfer agreements.\textsuperscript{17} Thus, both treatment and the control observations have MTAs and the difference between them is whether a license is issued.\textsuperscript{18} Table 3 presents the covariate balance for this sample.

*** Table 3 Here ***

Although the difference in means for the control and treatment groups in the MTA–linked sample is larger in absolute magnitude than that for the full sample reported in Table 2, the MTA sample reveals fewer statistically significant differences in means. For the MTA–linked sample, only the $t$–test of the difference of means for \textit{Publication Year} is highly significant (at the 1\% level), while the distributions between the treatment and control group observations are weakly significant (10\%) only for \textit{Journal Impact Factor}. This suggests a greater level of noise within the MTA–linked observations than in the full sample.

As we did in our analysis for the full sample, we present both the non–parametric citation curve for the matched samples in Figure 8 and the regression results in Figure 9 below.

*** Figures 8 and 9 Here ***

Our analysis of the effects of licensing on citations to patent–paper pairs that are associated with MTAs reveals a very different effect from that observed in the overall sample. For the overall sample, licensing is associated with an increase in citations to the publications linked to the underlying patented disclosure. For the MTA–linked sample, however, licensing is associated with a decline in citations. Using the covariate adjusted values (Figure 9), the magnitude of the licensing effect is -3.6 citations in year $t+1$ and -3.2 in year $t+3$ (and statistically significant at the 1\% level in those two years), but is not statistically significant from zero in year $t+2$.\textsuperscript{19} Overall, these coefficients represent a decrease in citations of approximately 30\% for the average publication, although our small sample size calls for caution in interpreting the exact magnitude of the effect.\textsuperscript{20}

The timing of the “license effect” for citations to MTA–linked patent–paper pairs also suggests a more rapid negative impact than was observed for the positive effect of licenses on citations in the overall sample.

\textsuperscript{17}We include MTAs that are issued before or after the license. In doing so we assume that an invention receiving an MTA irrespective of when the MTA was executed (i.e., the execution of the MTA) does not “convert” the material into a research tool. Restricting the sample to instances where an MTA exists prior to the license could ensure against possible reverse causality, however. This restriction led to results that are similar in direction, although with much smaller sample sizes. Thus we report results with the larger sample of MTA–linked inventions.

\textsuperscript{18}Because we also match treatment to control observations based on whether the observation has an MTA, treated observations with MTAs prior to license are matched to controls that also already have an MTA, while those that have not received an MTA prior to license are matched to controls that also have not yet received an MTA as of the license execution date.

\textsuperscript{19}We do not ascribe any economic meaning to the unusual result in $t+2$. This can be seen in robustness checks with 2–inventor overlap (coefficients of -5.3\*, -3.4\*, -5.1\*** for years $t+1$, $t+2$, and $t+3$) and 2–inventor overlap considering only patents with fewer than six publications (coefficients -5.9\***, -4.1\***, -5.9\***). Both show strong effects in $t+2$ as well as in $t+1$ and $t+3$.

\textsuperscript{20}For robustness, we also conducted our analysis of research tool licensing using the Murray and Stern (2007) patent class–based definition of research tools and obtained similar results.
The apparently more rapid appearance of the negative effects on citations observed in the MTA sample is broadly consistent with the delays and project abandonment observed in the surveys by Walsh et al. (2007) and Lei et al. (2009), as well as interviews we conducted for this study. Moreover, declines in citations for the MTA–linked disclosures that are licensed occur as early as the year of the execution of the license. Inasmuch as the citations to the focal publication associated with the patented disclosure appear only after a lag, the speed of this observed effect suggests that researchers or universities may limit or impose other restrictions on sharing of “research tools” during the negotiation of the license.\footnote{Note that the identification of any in–year effect is much weaker, since the exact timing of the license during the year is not accounted for in the analysis.}

6 Conclusion

This paper has investigated the effects of licenses on communication among scientists of published research results that are linked to patented academic invention disclosures within the University of California system. Our results suggest that in general, licenses on scientific work are associated with an increase in the number of citations to related publications, but that this effect differs for a class of patented invention disclosures that we believe includes a high share of research tools. For these inventions we observe a decrease in the number of citations following a license.

Our results are consistent with other findings in the literature that suggest that licensing may have a positive signaling effect, but that licenses on research tools may lead to restrictions on input materials that are important for follow–on research.

These findings should be interpreted with caution, however. Our citation data do not currently incorporate the identities of the individuals and organizations citing the scientific journals within our sample. Any “self–citations,” or citations made by the authors of the original work, would not represent knowledge flows. Their inclusion, therefore, could overstate the effect of licenses on scientific communication.

Accounting for citers’ identities would also allow us to investigate the extent to which follow–on research is conducted by licensees, thus suggesting the presence of market–based knowledge flows—or by third–parties, which would indicate spillovers operating within cumulative research. We are presently incorporating the identities of authors of citing journal articles into our data in order to address these issues in an upcoming revision.

This study makes three contributions. First, it introduces a method for linking patents and publications associated with an academic research advance, and thereby automates a previously arduous process that has been difficult to scale up. Second, it employs a matching methodology that avoids strong parametric assumptions that may not be appropriate for the complex interactions that underlie journal citation patterns.
And finally, but most importantly, this paper contributes to our understanding of the effect of the licensing of intellectual property rights covering academic research output on scientific communication.
References


National Science Foundation (2012). Science and Engineering Indicators.


Table 1: Summary Statistics for Samples with Three and Four Author Publication Matches to Patents

<table>
<thead>
<tr>
<th>Samples</th>
<th>3+</th>
<th>4+</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sample Size†</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Publications (000)</td>
<td>1.7</td>
<td>0.6</td>
</tr>
<tr>
<td>Patents (000)</td>
<td>0.7</td>
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</tr>
<tr>
<td>Publications with MTAs</td>
<td>261</td>
<td>79</td>
</tr>
<tr>
<td>Publications / Patent</td>
<td>2.4</td>
<td>1.8</td>
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<tr>
<td>Observations in the Life Sciences</td>
<td>49%</td>
<td>44%</td>
</tr>
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</table>

<table>
<thead>
<tr>
<th>Sample Statistics††</th>
<th></th>
<th>Difference†††</th>
</tr>
</thead>
<tbody>
<tr>
<td>Citations Per Year</td>
<td>11.4 (26.1)</td>
<td>16.2 (36.3)</td>
</tr>
<tr>
<td>Average Impact Factor</td>
<td>8.7 (8.4)</td>
<td>11.0 (9.8)</td>
</tr>
<tr>
<td>Publication Year</td>
<td>2000.7 (2.6)</td>
<td>2000.4 (2.8)</td>
</tr>
<tr>
<td>Publication Age</td>
<td>3.2 (2.6)</td>
<td>3.3 (2.7)</td>
</tr>
<tr>
<td>Age at MTA Issuance</td>
<td>2.4 (2.7)</td>
<td>2.6 (2.6)</td>
</tr>
<tr>
<td>Age at Patent Issuance</td>
<td>3.5 (2.0)</td>
<td>3.5 (2.0)</td>
</tr>
</tbody>
</table>

†Values in the parentheses are the number of standard deviations.
††Sample restricted to those publications with 7 years of citation data (only first 7 years of data included)
†††Comparison is a t-test between those with 3-inventor overlap (not 3+) and those with 4+.

Table 2: Covariate Balance for the Full Licensing Sample

<table>
<thead>
<tr>
<th>Covariates</th>
<th>Mean Treated</th>
<th>Mean Control</th>
<th>t-Test</th>
<th>KS Test</th>
</tr>
</thead>
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<td>Publication Age</td>
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<td>-</td>
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<td>MTA Issued</td>
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<tr>
<td>Journal Impact Factor</td>
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<td>Publication Year</td>
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<td>Citations in t-1</td>
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<td>6.1</td>
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<td>***</td>
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<td>-</td>
</tr>
<tr>
<td>Citation Slope from t-2 to t-1</td>
<td>2.3</td>
<td>1.9</td>
<td>***</td>
<td>***</td>
</tr>
</tbody>
</table>

Notes:
*** p<0.01  ** p<0.05  * p<0.10
†Journal Subject is a categorical variable, with each subject mapped to a random integer. Therefore the 4.3 listed has no literal meaning, but the equality between treatment and control, as well as well as the lack of any difference, reinforce the success of the exact matching.
Table 3: Covariate Balance for the Research Tools Sample

<table>
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<th></th>
<th></th>
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<td>MTA Issued</td>
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<td>-</td>
<td></td>
</tr>
<tr>
<td>Journal Impact Factor</td>
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<td>6.0</td>
<td>-</td>
<td>*</td>
<td></td>
</tr>
<tr>
<td>Publication Year</td>
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<td>2000.5</td>
<td>***</td>
<td>-</td>
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<td>Citations in t–1</td>
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<tr>
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<td>Citation Slope from t–2 to t–1</td>
<td>4.3</td>
<td>3.8</td>
<td>-</td>
<td>-</td>
<td></td>
</tr>
</tbody>
</table>

Note: *** p<0.01  ** p<0.05  * p<0.10

Figure 1: Academia Share of U.S. Basic Research Performance
Figure 2: Academia Share of U.S. Utility Patents

Figure 3: Sample Composition
1. Start with the patent

2. Find the inventors

Inventor(s): William Fenical, Paul Jensen, Xing Cheng

3. Look at the overlap in their publications

Figure 4: Inventor–Based Matching Example

Figure 5: Distribution of Patent–Paper Pairs with Two or More Names Common to Both Matched Patent and Published Paper
Figure 6: License Effect—Full Licensing Sample

Figure 7: Regression Results from the License Effect—Full Licensing Sample

(Error bars signify ± 1 standard error.)
Figure 8: License Effect—Research Tools

Figure 9: Regression Results from the License Effect—Research Tools

(Error bars signify ± 1 standard error.)