# On the Development and Distribution of R Packages: An Empirical Analysis of the R Ecosystem

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# ABSTRACT

This paper explores the ecosystem of software packages for R, one of the most popular environments for statistical computing today. We empirically study how R packages are developed and distributed on different repositories: CRAN, BioConductor, R-Forge and GitHub. We also explore the role and size of each repository, the inter-repository dependencies, and how these repositories grow over time. With this analysis, we provide a deeper insight into the extent and the evolution of the R package ecosystem.

# **Categories and Subject Descriptors**

D.2.7 [Software Engineering]: Distribution, Maintenance and Enhancement—*Version Control*; D.3.3 [Software Engineering]: Programming Languages

# Keywords

software ecosystem, software distribution, software development, package, R, software repository mining

# 1. INTRODUCTION

A wide range of environments for statistical computing have been proposed and used over the years. The most important ones are SAS, SPSS, R, Stata, Statistica, Minitab, Systat and JMP. While SAS and SPSS have been the market leaders in the past, R is catching up rapidly. According to Hornik [10], the number of R packages available in CRAN, the main and official distribution of R packages, is witnessing a slightly "sub-exponential" growth. Muenchen [12] counted the number of CRAN packages against each major release of R (until version 3.0.2), and observed a quadratic growth. He states: "To put this growth in perspective [...] During 2013 alone, R added more functions/procs than SAS Institute has written in its entire history!"

The R programming language also continues to increase in popularity. The Tiobe index for June  $2015^1$  ranked R as the 13th most popular programming language. The "Transparent Language Popularity Index"<sup>2</sup> ranked R as the 14th most popular programming language, all language categories confounded.

Because of this popularity of the R language and its software package ecosystem, it is important to study this ecosystem in more detail. In particular, we wish to get insight in the effect of other package repositories on CRAN.

Taking the point of view of a R package user, we focus on the following questions in this paper. Where and how are packages developed and distributed? How do packages depend on one another? Do we observe change over time related to these questions?

The R community has raised concerns about the way R packages are currently distributed, what problems current package management systems suffer from, and how they could be solved [13]. Among others, we wish to know whether the increasing popularity of GitHub as a host for developing many R packages has become a "game changer". Indeed, it is quite straightforward to develop and install packages directly from GitHub, possibly avoiding the need for having one's package distributed on CRAN or BioConductor.

The remainder of this paper is structured as follows. Section 2 discusses the R package repositories that will be considered as part of our analysis. Section 3 presents related work. Section 4 explains how we have extracted all data necessary for our analysis. Section 5 presents the results of our exploratory empirical analysis. Section 6 discusses the threats to validity of our research, Section 7 presents future work, and Section 8 concludes.

#### 2. CONSIDERED R REPOSITORIES

The R ecosystem is based on software packages. The main R distribution installs by default a dozen or so base packages, and another dozen or so recommended packages (the exact number depends on the installed version of R). These packages will be excluded from the analysis in this paper. In addition to these core R packages, many additional packages are developed and distributed by different contributors through different repositories.

# 2.1 R package distributions

<sup>1</sup>www.tiobe.com/index.php/content/paperinfo/tpci <sup>2</sup>lang-index.sourceforge.net, July 2013 update.

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Repository	URL	Number of pack-	Role	Development status	
name		ages [ Date ]			
CRAN	cran.r-project.org	6411 [19-03-2015]	Distribution only	Stable package releases only	
BioConductor	bioconductor.org	997 [19-03-2015]	Distribution only	Stable package releases only	
GitHub	github.com	5150 [17-02-2015]	Mainly development,	Git version control	
			rolling release distro		
			(e.g., with devtools)		
R-Forge	r-forge.r-project.org	1883 [18-03-2015]	Mainly development,	SVN version control	
			rolling release distro		
			(e.g., with devtools)		

 Table 1: Characteristics of considered R package repositories

CRAN, the Comprehensive R Archive Network, can be considered as the official repository that contains the broadest collection of R packages. It aims at providing stable and high-quality packages compatible with the latest version of R. Quality is ensured by forcing package maintainers to follow a strict policy. Packages are tested daily using the command-line tool R CMD check. When these tests fail, package maintainers have to resolve the problems before the next major R release. Problematic packages are archived, making it impossible to install them automatically<sup>3</sup>, as they will no longer be included in CRAN until a new package version is released that resolves the problems.

Another important distribution of R packages is BioConductor (abbreviated to BioC hereafter), focusing on software packages and datasets dedicated to bioinformatics. The datasets are subdivided into experiment data and annotation data. Together, they make up more than half of all packages available on BioC. Since the focus of this paper is on R's *software* ecosystem, these datasets will be treated separately during our analysis.

While fully compatible with the format of CRAN packages, BioC packages are not installed by default: users must configure their R installation with a BioC mirror. As for CRAN, packages that fail the daily check will be dropped from the next release of  $BioC.^4$ 

#### 2.2 R package development forges

While CRAN and BioC are the largest and most wellknown package repositories for the *distribution* of R packages, other repositories are frequently used for the *development* of R packages. Thanks to R packages like **devtools** (available since 2011), development of R packages is facilitated, and direct installation of packages from a source code repository like Git or Subversion becomes straightforward. As such, there is no longer a strict need to rely on package distributions. Therefore, R development forges have become an important and integral part of the R package ecosystem.

The two R package development forges that we consider in our study are *R*-Forge and GitHub. *R*-Forge is specialized at hosting R code. Its main target is to provide a central platform for the development of R packages, offering SVN repositories, daily built and checked packages, bug tracking, and so on. GitHub, a web platform for Git's distributed version control system, is also becoming increasingly popular

www.bioconductor.org/checkResults

for R package development.

Table 1 provides a brief comparison of the four R package repositories considered in our study. It also provides an idea of the size of each repository, in terms of the number of provided R packages. When installing packages from CRAN or BioC, only the most recent version is guaranteed to work with the latest version of R.

Support for multiple repositories is built deeply into R. For example, the R function install.packages can take the source repository as an optional argument, or can be used to install older versions of a given package. While *R*-Forge and *GitHub* are focused on development, packages can be installed directly from these forges using functions of the devtools package. For example, the function install\_github allows the installation of R packages directly from *GitHub*, while the function install-svn allows the installation from a Subversion repository (such as the one used by *R*-Forge). By default, these functions will install the latest package version, but optional parameters can be used to install a specific commit.

#### **3. RELATED WORK**

Our research aims at studying the characteristics of R packages in function of the repository in which they are developed or distributed. A similar question, though not restricted to R packages, was studied by Beecher et al. in [3, 1]. They explored how the structure, complexity and decay of open source projects may be influenced by the repository in which they are retained (e.g., SourceForge, Savannah, Debian, RubyForge, GNOME, KDE). They concluded that membership of a particular repository may depend on the maturity and quality of the project. For example, SourceForge tends to host more early inceptors and immature projects, while Debian tends to hosts high-quality mature projects.

A number of researchers have studied the evolution of R packages. We are not aware of any studies taking into account other R package distributions (such as BioC) or development forges (such as *R*-Forge or GitHub). All related research seems to be restricted to the official package distribution CRAN.

German et al. [7] studied the evolution of CRAN from an ecosystemic viewpoint. They compared the characteristics, growth, dependencies and community structure of core packages and user-contributed packages. They also analysed the user and develop communities by studying mailing list traffic. In earlier work [5], we have studied the maintainability of CRAN packages in terms of errors discovered by the R CMD check tool, and how this relates to package depen-

 $<sup>^{3}\</sup>mathrm{It}$  is still possible to install them manually.

<sup>&</sup>lt;sup>4</sup>The check results and available packages for current and past releases of BioC are available on

dencies and package updates. Based on these insights, in [4] we presented a web-based dashboard for helping R package maintainers deal with such issues.

There has been quite some empirical research on the software ecosystem of *GitHub* repositories, though not specifically related to R packages. Dabbish et al. [6] focused on the social and community aspects of *GitHub*. Vasilescu et al. [14] compared the involvement on *GitHub* with the activity on Stack Overflow, a Q&A website for software developers. Vasilescu et al. [15] studied a large sample of *GitHub* projects developed in Java, Python and Ruby. They compared direct code modifications (commits) with indirect ones (pull requests) and related this to success or failure of continuous integration with TRAVIS-CI. Gousios et al. [8, 9] provide *GHTorrent*, a scalable way to analyse *GitHub* data.

#### 4. DATA EXTRACTION

An R package is a tar.gz archive that can contain code, examples, tests, data sets, and so on. It also includes a mandatory *DESCRIPTION* file that contains metadata describing the package's characteristics such as name, purpose, maintainer, version, imports and dependencies.

In order to carry out our empirical analysis, we need to extract data from each of the targeted R package repositories. We wrote a set of R and Python scripts to retrieve and process the data. A replication package is available on github.com/ecos-umons/iwseco2015.

**CRAN** – The metadata of R packages on *CRAN* was retrieved using  $extractoR^5$ , an R package that we already developed and used in earlier work [5]. It downloads the *CRAN* package sources, extracts their contents and stores the *DESCRIPTION* file metadata into R *data.frame* objects. Using this tool we collected, since September 2013, the metadata of 7,654 packages with their associated versions and dependencies. We explored in detail the state of *CRAN* on 19 March 2015, containing 6,411 non-archived packages.

BioC – On the same date, we extracted the metadata from the *DESCRIPTION* files of all 997 R software packages corresponding to release 3.0 of *BioC*. A list of candidate packages was retrieved from the SVN<sup>6</sup> and the *DE-SCRIPTION* files were retrieved. We separately considered *annotation data* packages and *experiment data* packages for our analysis.

R-Forge – We first identified 1,883 repositories on R-Forge on 18 March 2015 using the full repository list<sup>7</sup>. Some of those repositories contained no package, while others contained more than one package, since we identified several DESCRIPTION files corresponding to distinct package names. An automatic script extracted the metadata of 2,217 packages based on the DESCRIPTION files contained in those 1,883 repositories.

GitHub – GitHub Archive<sup>8</sup> is a tool that collects and stores data from GitHub that usually are not available anymore on GitHub after some time. We relied on the events stream collected by GitHub Archive to identify potential active R repositories on GitHub. First, we collected from

GitHub Archive all events that occurred for two years until 31 December 2014, when GitHub released its new (partially backward incompatible) API. Then, we filtered out all events of type *PushEvent* that were related to repositories whose language was flagged 'R' by their owner. This resulted in a list of 121,385 candidate repositories that could contain R packages. We checked for the presence of a DESCRIP-TION file at the root of those repositories, and fetched this file on 17 February 2015. As we are interested in comparing GitHub with other package sources like CRAN and BioC, if packages with the same name were found in two or more distinct repositories (including forks), we assumed the package belonging to the oldest repository to be the main one. Importantly, we also filtered out the repositories belonging to the *GitHub* accounts cran and rpkg, as these two accounts were used to mirror (part of) the contents of CRAN. This resulted in 5,150 distinct R packages from *GitHub*.

#### 5. EMPIRICAL RESULTS

# 5.1 Where and how are R packages developed and distributed?

Figure 1 shows the overlap of R packages on different distributions and development forges. The overlap between CRAN and BioC is very limited, as expected. Both package distributions only have 4 packages in common, corresponding to 0.4% of all considered BioC packages and only 0.06% of all considered CRAN packages.

Many packages on CRAN also appear on GitHub, since both repositories serve different purposes (distribution and development, respectively). We also observe in Figure 1 that the intersection of packages that can be found on CRAN and GitHub is non negligible. 18.1% of all considered CRANpackages can also be found on GitHub. 22.5% of all R packages on GitHub are also present on CRAN. This relatively large proportion of overlap can be explained by the fact that CRAN is only a distribution platform, and cannot be used for collaborative development. Hence, **many R packages are developed on GitHub, while stable releases of these packages are published on CRAN**. We observe something similar when comparing BioC with GitHub, and for the same reasons. Indeed, 20.6% of all packages on BioChave a counterpart on GitHub.

*R-Forge* has 12.3% of its packages in common with *GitHub*, while as much as 45.2% of its packages are in common with *CRAN*. This shows that *R-Forge* serves as a development platform for *some* of the packages that get distributed through *CRAN*. This is not true for *BioC*: only 1.1% of all *R-Forge* packages are also available on *BioC*.

One of our goals is to study whether development forges like GitHub are overtaking CRAN and BioC as a primary source of R packages. To do so, we consider the set of all R packages that are available on GitHub on 17 February 2015, and study since when they were created and had a counterpart in other R package repositories.

Figure 2 suggests that the monthly number of newly created repositories for CRAN and BioC packages on GitHubis slightly increasing over time. This seems to imply that, over time, **developers of packages that are distributed on** CRAN and BioC decide to use GitHub as a host for developing their packages. This does not seem to affect the growth of the packages in the CRAN and BioCdistributions. Figure 3 shows that the number of packages in

<sup>&</sup>lt;sup>5</sup>github.com/ecos-umons/extractoR

<sup>&</sup>lt;sup>6</sup>hedgehog.fhcrc.org/bioconductor/branches/RELEASE\_3\_ 0/madman/Rpacks/ (username/password: readonly)

<sup>&</sup>lt;sup>7</sup>r-forge.r-project.org/softwaremap/full\_list.php

<sup>&</sup>lt;sup>8</sup>www.githubarchive.org

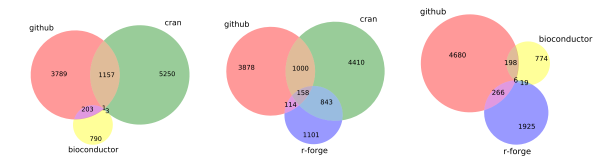


Figure 1: Intersections of R packages belonging to GitHub, CRAN, BioC and R-Forge

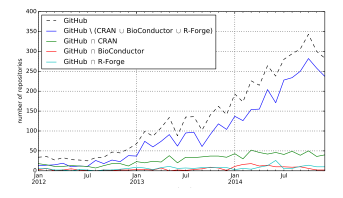


Figure 2: Monthly number of newly created repositories on *GitHub* containing R packages.

 $GitHub \cap CRAN$  grows faster than the number of packages distributed on CRAN.

We did not find evidence of packages disappearing from CRAN or BioC due to their migration to GitHub. As shown in Figure 1, in March 2015 CRAN and BioC still remain the primary sources for the *distribution* of stable R packages. As such, **development of R packages through** GitHub seems to complement distribution of packages through CRAN and BioC, and perhaps even has a catalyst effect.

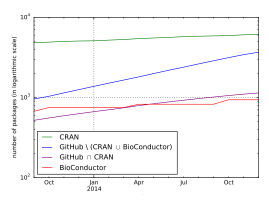


Figure 3: Evolution of the number of R packages in CRAN, GitHub and BioC.

Figures 2 and 3 also reveal that GitHub is increasingly hosting R packages that do not have a counterpart

in *CRAN* or *BioC*. Many of these packages are no longer actively maintained today. Those that do, may be developed for personal use only, or could still be unstable but at some point in the future may turn into stable packages that could become distributed in *CRAN* or *BioC*.

#### 5.2 How do packages depend on one another?

Every R package needs to specify in its *DESCRIPTION* file the packages it depends upon. In our analysis we consider as dependencies those packages that are listed in the **Depends**: and **Imports**: fields of the *DESCRIPTION* file, as these are the ones that are required to install and load a package.<sup>9</sup>

If an R package depends on another, do these packages belong to the same repositories, or do we observe many interrepository dependencies? We expect most of such interrepository dependencies to go towards CRAN since it is the official R package distribution. We also expect many dependencies from other repositories towards BioC since it is an active package distribution that offers the same quality checks as CRAN, and also because it contains many useful datasets. Therefore, when considering dependencies to packages belonging to BioC, we also count dependencies to dataset packages belonging to BioC.

As can be seen in Figure 1, the same package may belong to different repositories (for example, GitHub may store the development version while CRAN may contain the stable release version of the package). To cope with this, we define a total order > on the set  $S = \{CRAN, BioC \text{ software}, BioC$ datasets, GitHub, R-Forge, Unknown $\}$  such that CRAN >BioC software > BioC datasets > GitHub > R-Forge > Unknown. This total order privileges the distributed version of a package over its development version. For example, if a package  $p_1$  on GitHub depends on a package  $p_2$  that belongs to both CRAN and GitHub, it will be counted as a dependency from GitHub to CRAN.

Using this total order we can introduce some dependency metrics. Let  $\alpha, \beta \in S$  be two R package repositories (i.e., sets of packages). We define:

$$deps(\alpha, \beta) = \{ (p_1, p_2) \in \alpha \times \beta \mid \\ p_1 \text{ depends on } p_2 \land \nexists \gamma \in \mathcal{S} : (p_2 \in \gamma \land \gamma > \beta) \}$$
$$|\{p_1 \in \alpha \mid \exists p_2 \in \beta : (p_1, p_2) \in deps(\alpha, \beta) \}$$

dependsOn
$$(\alpha, \beta) = \frac{|\{p_1 \in \alpha \mid \exists p_2 \in \beta : (p_1, p_2) \in deps(\alpha, \beta)\}|}{|\alpha|}$$

$$\operatorname{requiredBy}(\alpha,\beta) = \frac{|\{p_1 \in \alpha \mid \exists p_2 \in \beta : (p_2,p_1) \in \operatorname{deps}(\beta,\alpha)\}|}{|\alpha|}$$

<sup>9</sup>R CMD check also verifies Suggests: dependencies.

<u> </u>								
		β						
Metrics	$\alpha$	CRAN	BioC software	BioC datasets	GitHub	<i>R</i> -Forge	Unknown	
dependsOn $(\alpha, \beta)$		61.0%	2.1%	0.1%	0%	0%	0%	
requiredBy $(\alpha, \beta)$	CRAN	24.9%	5.8%	0.1%	15.7%	8.6%	_	
$ deps(\alpha,\beta) $		10,560	212	4	1	1	0	
dependsOn $(\alpha, \beta)$	BioC	58.8%	77.1%	9.3%	0.2%	0.1%	1.1%	
requiredBy $(\alpha, \beta)$	software	6.5%	$\mathbf{26.5\%}$	2.8%	12.6%	4.8%	_	
$ deps(\alpha,\beta) $		$1,\!615$	2,748	133	2	1	13	
dependsOn $(\alpha, \beta)$	BioC	15.2%	77.1%	17.5%	0%	0%	0%	
requiredBy $(\alpha, \beta)$	datasets	0.2%	6.2%	4.3%	1.9%	0.3%	_	
$ deps(\alpha,\beta) $	(1,115 packages)	333	$1,\!567$	204	0	0	0	
dependsOn $(\alpha, \beta)$		48.9%	5.2%	7.4%	5.7%	0.3%	2.7%	
requiredBy $(\alpha, \beta)$	GitHub	0%	0%	0%	4.9%	0.1%	-	
$ deps(\alpha,\beta) $		8,614	684	37	386	15	156	
dependsOn $(\alpha, \beta)$		37.2%	2.3%	0.1%	0.7%	5.8%	1.5%	
requiredBy $(\alpha, \beta)$	R-Forge	0.1%	0.1%	0%	0.6%	5.0%	_	
$ deps(\alpha,\beta) $		1,830	93	4	19	136	36	

Table 2: Number of packages primarily belonging to repository  $\alpha$  that depend on or are needed by at least one package primarily belonging to repository  $\beta$ .

 $|\text{deps}(\alpha, \beta)|$  counts all dependency relationships from packages in  $\alpha$  to packages in  $\beta$ , dependsOn $(\alpha, \beta)$  gives the ratio of distinct packages in  $\alpha$  depending on at least one package in  $\beta$ , and requiredBy $(\alpha, \beta)$  the ratio of distinct packages in  $\alpha$  on which at least one package in  $\beta$  depends. Table 2 presents these metrics for all pairs of considered R package repositories. Unknown represents those dependencies for which we did not find a matching package name in any of the considered repositories. This value was especially high for *GitHub* (140 packages with 156 dependencies to 89 unknown packages).

CRAN is self-contained: the majority of dependencies of its packages stay within CRAN: 61% of all CRAN packages depend on another CRAN package. This is expected, since otherwise the packages would not pass the R CMD check. Note that only 24.9% of all CRAN packages are required by other CRAN packages.

**BioC** depends primarily on *CRAN* and on itself: 58.8% of all *BioC* packages depend on *CRAN* packages, while 77.1% of all *BioC* packages depend on other *BioC* packages. Similar to *CRAN*, 26.5% of all *BioC* packages are required by other *CRAN* packages. We also observe that 9.3% of *BioC* software packages depend on *BioC* datasets.

*GitHub* and *R-Forge* depend primarily on *CRAN*: 87.1% of *GitHub* dependencies and 86.4% of *R-Forge* dependencies go to *CRAN* packages.

This shows that **CRAN** is still at the center of the **ecosystem** and that it has a minority of packages forming a core required by other packages both from CRAN and other sources.

# 6. THREATS TO VALIDITY

We only considered a subset of the R ecosystem, consisting of only 4 package repositories, but covering more than 12,000 distinct R software packages. While other R package repositories exist, given their small size we have not included them in our analysis. The Omega Project for Statistical Computing" (www.omegehat.org) hosts around one hundred R packages. *RForge* (rforge.net), not to be confused with *R-Forge*, provides a collaborative environment for R package developers based on SVN repositories, and contains less than two hundred packages, many of which are no longer active. *GitHub* competitors like *BitBucket* (bitbucket.org), *Gitorious* (www.gitorious.com) and *Gitlab* (gitlab.com) are considerably less frequently used for hosting R package development.

While for *BioC* we explicitly excluded (or treated differently) the packages containing datasets, we were not able to do the same for the other repositories, since we found no automated way to distinguish "ordinary" software packages from datasets. If an R package contains both data and functions, it is hard to decide whether it should be regarded as a software package or a dataset.

For part of our analysis, we relied on information extracted from SVN or Git, or from hosting services like GitHub. There are many potential perils and pitfalls that should be taken into consideration when doing so [2, 11]. Some of them can be avoided, others or inherent to the limitations of the considered version control systems or hosting services. For example, how should forking be taken into account? In our analysis, we excluded all forks. We also relied on *GithubArchive* as a proxy of *GitHub* data to extract events, but we cannot guarantee that this data is fully consistent and complete.

For GitHub, we assumed that the R package DESCRIP-TION file always resides in the root directory of each Git repository. While this may have lead to the exclusion of some packages in our analysis, it avoids the inclusion of many "false positive" packages that are identical clones of existing CRAN package sources used by developers to avoid external dependency problems.

We also found that some GitHub accounts containing R packages (in particular, accounts cran and rpkg) actually served as partial mirrors of CRAN. These accounts were excluded from our analysis, but we have no guarantee that other accounts may also be mirrors of R packages developed or distributed elsewhere.

The chosen date of the R package ecosystem snapshot, and the chosen duration for the historical analysis may influence our results. Repeating the same analysis for other dates would allow us to confirm the observed results. For the historical analysis of the GitHub data, we based ourselves on the packages still existing in GitHub in February 2015. We were not able to extract historical information from GitHub repositories that have been removed before that date.

### 7. FUTURE WORK

Our analysis of the R software package ecosystem can be extended in many ways. An important avenue of future research is to study the evolution over time: do we observe important changes in the *diversity* of the ecosystem over time? Is there a relationship between the *quality* and *survival* of the packages contained in each considered repository? How does the *popularity* of packages (e.g., expressed in number of downloads or number of incoming dependencies) relate to these characteristics? How easy is it to *install* R packages depending on their provenance? What happens to R packages (especially in *CRAN* and *BioC*) after they have become *archived*?

Another future research track concerns a *socio-technical analysis* of the R package ecosystem: how do package developers collaborate? Do we observe a different collaboration behaviour depending on the considered repository? Does this behaviour correlate with desirable properties of packages?

Finally, we wish to study the impact of ongoing initiatives to facilitate R package management and producing reproducible results. Examples are the *Drat R Archive Template*<sup>10</sup>, the *Managed R Archive Network*<sup>11</sup>, the *Reproducible R Toolkit*<sup>12</sup>, and packrat, a portable and reproducible dependency management system for R projects.

#### 8. CONCLUSION

In this paper, we studied the ecosystem of R packages beyond the official CRAN repository. We also considered the *BioConductor* package distributions, and we explored two R package development forges: *GitHub* and *R-Forge*. In total, we analysed the origin and the dependencies of more than 12,000 packages that were still available in March 2015.

We observed that CRAN remains the center of the R package ecosystem, since its packages do not depend on external packages, while *BioConductor*, *R-Forge* and *GitHub* strongly depend on *CRAN* packages. *BioConductor* also contains many packages required by the others, but with an order of magnitude difference compared to *CRAN*.

We also observed that GitHub is becoming increasingly used as a collaborative development platform for R packages, both for packages already distributed on CRAN and BioConductor, as well as for new packages that do not have any counterpart in the considered distributions or forges. We did not observe any positive or negative effect of this increased use of GitHub on the growth of the number of CRANor BioConductor packages.

In the future we intend to explore "the *GitHub* effect" in more detail, taking into account other factors such as the quality and longevity of packages, as well as a possible effect on the collaborative behaviour of package developers.

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