

FORESEE: a tool for the systematic comparison of translational drug response modeling pipelines

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Abstract

Translational models that utilize *omics* data generated in *in vitro* studies to predict the drug efficacy of anti-cancer compounds in patients are highly distinct, which complicates the benchmarking process for new computational approaches. In reaction to this, we introduce the uniFied translatiOnal dRug rESponse prEdiction platform FORESEE, an open-source R-package. FORESEE not only provides a uniform data format for public cell line and patient data sets, but also establishes a standardized environment for drug response prediction pipelines, incorporating various state-of-the-art preprocessing methods, model training algorithms and validation techniques. The modular implementation of individual elements of the pipeline facilitates a straightforward development of combinatorial models, which can be used to re-evaluate and improve already existing pipelines as well as to develop new ones.

Availability and Implementation. FORESEE is licensed under GNU General Public License v3.0 and available at <https://github.com/JRC-COMBINE/FORESEE>.

Supplementary Information. Supplementary Files 1 and 2 provide detailed descriptions of the pipeline and the data preparation process, while Supplementary File 3 presents basic use cases of the package.

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33 1 Introduction

34 Cell line data bases featuring both multi-omics characterizations of human cancer cell lines and their
35 response profiles to drug compounds have become a vital tool in developing predictive drug response
36 models for cancer patients. Concomitant with their advancement, tools have been developed to provide
37 access to the data (Smirnov et al., 2015; Luna et al., 2016) and to systematically evaluate models for drug
38 response prediction (Jang et al., 2014). In order to attain clinical relevance, such cell line-based models
39 need to be translated and tested on patient data, which has become the focus of a steadily growing
40 number of studies: while some are restricted to gene expression data (Geeleher et al., 2014; Huang et al.,
41 2017), other studies additionally incorporate mutation profiles and copy number variations (Dorman
42 et al., 2016), promoter methylation (Aben et al., 2016) and protein expression (Daemen et al., 2013).
43 As a consequence of the diversity and scope of the work that has been performed in this field, comparing
44 the various approaches is complicated and the process of benchmarking novel computational methods
45 has become time-consuming. In order to address this, we introduce the FORESEE platform to facilitate
46 a straightforward and comprehensive evaluation of translational drug response models.

47 2 Implementation

48 For the systematic evaluation of individual components of the modeling pipeline and their impact on
49 the performance, the FORESEE package features not only functional elements of the pipeline, but also
50 introduces a common data format for frequently used data resources. Thus, it allows for the methodical
51 investigation of all possible combinations of modeling choices, as well as for testing a specific pipeline
52 on different data sets, thereby exploring how the choice of data affects modeling performance.

53 2.1 Data

54 Supporting the idea of translational modeling pipelines, the FORESEE package comprises molecular
55 and pharmacological data that characterize cell lines, xenografts and patients. In terms of cell line
56 characterization, data from the GDSC (Garnett et al., 2012), the CCLE (Barretina et al., 2012; Cancer
57 Cell Line Encyclopedia Consortium and Genomics of Drug Sensitivity in Cancer Consortium., 2015), the
58 CTRP (Rees et al., 2016; Seashore-Ludlow et al., 2015; Basu et al., 2013) and Daemen et al. (Daemen
59 et al., 2013) were formatted into *ForeseeCell* objects. Each of these *ForeseeCell* objects contain at least
60 one type of molecular data, such as gene expression, and one type of pharmacological data, such as the
61 IC_{50} (half maximal inhibitory concentration). On the other hand, information of patients with breast
62 cancer (GSE6434 (Chang et al., 2005) and GSE18864 (Silver et al., 2010)), lung cancer (GSE33072 (Byers
63 et al., 2013)), ovarian cancer (GSE51373 (Koti et al., 2013)) and multiple myeloma (GSE9782 (Mulligan
64 et al., 2007)) was organized into *ForeseePatient* objects including at least one molecular data type and
65 one measure of *in vivo* drug efficacy, such as tumor shrinkage. Although primarily developed to design
66 translational models that are trained on cell line data and tested on patient data, FORESEE also allows
67 for testing cell line-trained models on other cell line data sets. Moreover, data from patient derived
68 xenografts (PDX) (Gao et al., 2015; Witkiewicz et al., 2016), bridging the differences between cell
69 lines and patients, was included as *ForeseeCell* objects to offer a supplementary translational modeling
70 opportunity: testing PDX-trained models on patients. A detailed description of how the data was
71 obtained and prepared can be found in Supplementary File 2.

72 2.2 Pipeline

73 The functional elements of the modeling pipeline, which are depicted in Figure 1 and explained in
74 more detail in Supplementary File 1, are implemented as independent modules that can be changed
75 individually, according to the user's preferences. Across all main steps of the pipeline, user-defined
76 functions can substitute the pre-implemented methods to enable a more flexible use of the package,
77 which is explained in Supplementary File 3 along with other use cases.

78 3 Discussion

79 The FORESEE R package is designed to explore and compare translational drug response models. Thus,
80 it comprises both a standardized data format for molecular *in vitro* and *in vivo* data and functional
81 building blocks that summarize various well-established preprocessing and processing options. Moreover,
82 each of the functional blocks allows for the application of user-defined alternatives to support the fast and
83 easy development of novel modeling pipelines. Future expansions of FORESEE are directed towards an
84 automatic optimization for identifying the modeling pipeline best-suited for a particular setting. Until
85 then, we hope that FORESEE can facilitate exchanging expertise among researchers by providing a
86 standard environment for translational drug sensitivity models and therefore push forward the potential
87 to predict drug sensitivity of cancer patients.

88 4 Availability

89 The FORESEE package is available at <https://github.com/JRC-COMBINE/FORESEE> and
90 <https://doi.org/10.17605/OSF.IO/RF6QK>, and provides vignettes for documentation and application
91 both online and in the Supplementary Files 2 and 3.

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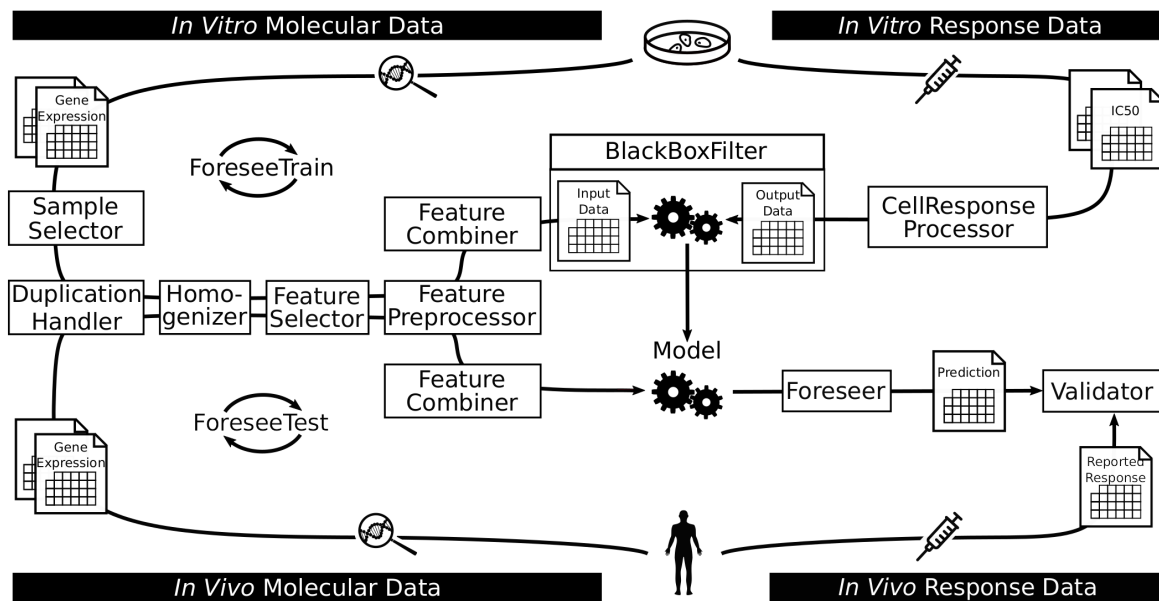


Figure 1: Illustration of the general FORESEE pipeline. The modeling routine comprises two main shells, ForeseeTrain (upper loop) and ForeseeTest (lower loop), with each consisting of different functional elements (boxes). During training, molecular cell line data is preprocessed by selecting certain samples in SampleSelector, removing duplicated feature names in DuplicationHandler, reducing batch effects in Homogenizer, selecting certain features in FeatureSelector, transforming the data in FeaturePreprocessor and combining the different molecular data types in FeatureCombiner, while the response data is transformed in CellResponseProcessor. The preprocessed data is then used for model training in BlackBoxFilter. The Foreseer applies the completed model to molecular patient data that has been preprocessed in the same manner as the cell line data to yield a prediction for patient drug sensitivity, which is subsequently compared to the actual response in Validator to evaluate the overall performance of the translational model.

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