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# Automated residual plot assessment with the R package autovi and the Shiny application autovi.web

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### Summary

Visual assessment of residual plots is a common approach for diagnosing linear models, but it relies on manual evaluation, which does not scale well and can lead to inconsistent decisions across analysts. The lineup protocol, which embeds the observed plot among null plots, can reduce subjectivity but requires even more human effort. In today's data-driven world, such tasks are well-suited for automation. We present a new R package that uses a computer vision model to automate the evaluation of residual plots. An accompanying Shiny application is provided for ease of use. Given a sample of residuals, the model predicts a visual signal strength (VSS) and offers supporting information to help analysts assess model fit.

Key words: initial data analysis; statistical graphics; data visualisation; visual inference; computer vision; machine learning; hypothesis testing; regression analysis; model diagnostics

#### 1. Introduction

Regression analysis is a widely used statistical modelling technique for data in many fields. There is a vast array of software for conducting regression modelling and generating diagnostics. The package lmtest (Zeileis & Hothorn 2002) provides a suite of conventional tests, while the stats package (R Core Team 2022) offers standard diagnostic plots such as residuals vs. fitted values, quantile-quantile (Q-Q) plots, and residuals vs. leverage plots. Additional packages like itools (Long 2022), olsrr (Hebbali

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2024), rockchalk (Johnson 2022), and ggResidpanel (Goode & Rey 2019) deliver similar 16 graphical diagnostics, often with enhanced aesthetics or interactive features. These 17 tools collectively produce the core diagnostic plots outlined in the classical text by Cook 18 & Weisberg (1982). The ecostats package (Warton 2023) extends these diagnostics by 19 incorporating simulation envelopes into residual plots. Meanwhile, DHARMa (Hartig 20 2022) compares empirical quantiles (0.25, 0.5, and 0.75) of scaled residuals to their 21 theoretical counterparts, with a strong focus on identifying model violations such as 22 heteroscedasticity, misspecified functional forms, and issues specific to generalised 23 linear and mixed-effect models, like over/under-dispersion. It also provides conventional 24 test annotations to reduce the risk of misinterpretation. 25

However, relying solely on subjective assessments of these plots can lead to issues such as 26 over-interpreting random patterns as model violations. Li et al. (2024a) demonstrated 27 that visual inference methods, particularly those using the lineup protocol (Buja 28 et al. 2009), offer more practical and reliable assessments of residual patterns than 29 conventional tests, as they are less sensitive to minor departures. Packages such as 30 nullabor (Wickham et al. 2020), HLMdiag (Loy & Hofmann 2014), and regressinator 31 (Reinhart 2024) support this approach by enabling users to compare observed residual 32 plots with plots generated under null hypothesis, thereby helping to quantify the 33 significance of any detected patterns. 34

As noted in Li et al. (2024b), the lineup protocol has significant limitations in large-35 scale applications due to its reliance on human labour. To overcome this constraint, 36 a computer vision model was developed alongside a corresponding statistical testing 37 procedure to automate the assessment of residual plots. The model takes as input 38 a residual plot and a set of auxiliary variables (such as the number of observations) 39 and outputs a predicted visual signal strength (VSS). This VSS estimates the degree 40 of deviation between the residual distribution of the fitted model and the reference 41 distribution expected under correct model specification. 42

To make the statistical testing procedure and trained computer vision model widely accessible, we developed the R package autovi along with a companion web interface, autovi.web, which allows users to automatically assess their residual plots using the trained computer vision model.

The remainder of this paper is structured as follows: Section 2 introduces the definition and computation of visual signal strength. Section 3 expands on the computation of the null and bootstrapped residuals. Section 4 provides a detailed documentation of the autovi package, including its usage and infrastructure. Section 5 focuses on the

- 51 autovi.web interface, describing its design and usage, along with illustrative examples.
- 52 Finally, Section 6 presents the main conclusions of this work.

# 2. Definition and computation of visual signal strength

- 54 To train a computer vision model, a measure of the visible pattern in a plot is needed.
- 55 We call this the visual signal strength (VSS), which measures how prominently a
- specific set of visual patterns appears in an image. This can be computed for a training
- set of data, and plots, where the generating distributions are specified.
- 58 In the context of classical normal linear regression model diagnostics, VSS describes
- 59 the clarity of visual patterns on a diagnostic plot that may indicate model violations.
- 60 Violations can be categorised as weak, moderate, or strong, but here we treat it
- as a continuous positive real variable. Importantly, its interpretation depends on
- 62 how it is linked to a function of the data or the underlying data generating process.
- 63 Consequently, the calculation of VSS can vary across different model classes or within
- the same model, depending on the generating function.
- 65 VSS estimates the distance between the residual distribution of a fitted classical normal
- 66 linear regression model and a reference distribution (see Li et al. 2024b, for details).
- 67 The distance measure is based on the Kullback-Leibler (KL) divergence:

$$D = \log \left(1 + D_{KL}\right).$$

where  $D_{KL}$  is given by:

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$$D_{KL} = \int_{\mathbb{P}^n} \log \frac{p(\hat{e})}{q(\hat{e})} p(\hat{e}) d\hat{e}, \tag{1}$$

- here,  $\hat{e}$  denotes the residual vector from the regression model, and  $p(\cdot)$  and  $q(\cdot)$  are
- 70 the probability density functions of the reference residual distribution P and the true
- 71 residual distribution Q, respectively.
- 72 This distance measure depends on knowledge of the true residual distribution, which
- 73 is unknown in practice. To compute  $D_{KL}$  for the training samples, Equation (1)
- 74 takes different forms depending on the specific model violations. For instance, where
- 75 necessary higher-order predictors, Z, and their corresponding parameter,  $\beta_Z$ , are
- 76 omitted from the fitted linear model, the distance measure can be expanded as follows:

$$D_{KL} = \frac{1}{2} \left( \boldsymbol{\mu}_z^\top (\mathrm{diag}(\boldsymbol{R} \sigma^2))^{-1} \boldsymbol{\mu}_z \right),$$

where  $\mu_z = RZ\beta_z$ ,  $R = I_n - X(X^\top X)^{-1}X^\top$  and X is the design matrix of the regression model.

The computer vision model approximates this mapping from a set of residuals to 79 its corresponding distance measure. It is trained on a large number of synthetic 80 regression models, each designed to simulate specific violations of classical linear 81 regression assumptions. These models incorporate non-linearity through Hermite 82 polynomial transformations of predictors, heteroscedasticity by making the error 83 variance a predictor-dependent function, and non-normality by drawing residuals from 84 distributions such as discrete, uniform, and lognormal. Both simple and multiple linear 85 regression structures are used, with controlled parameters to generate diverse and 86 complex residual patterns. Since the data-generating process is known, the distance measure D can be explicitly calculated, enabling supervised training. The computer 88 vision model takes a residual plot as input and outputs the corresponding distance 89 measure, learning to quantify model violations directly from visual patterns. Additional 90 details are provided in Li et al. (2024b). 91

## 3. Definition and simulation of null and bootstrapped residuals

In the subsequent sections, we will frequently refer to null residuals and bootstrapped residuals, so it is helpful to first define and explain how they are generated.

Null residuals are used to generate null plots within the lineup protocol framework, 95 serving as the foundation for the statistical testing in our automated residual plot 96 assessment. Specifically, they represent residuals generated under the null hypothesis 97 that the model is correctly specified. A common method for simulating null residuals 98 in linear regression involves sampling from a normal distribution with mean zero and 99 variance equal to the estimated variance of the error term. These simulated residuals 100 and their corresponding plots depict what one would expect from a correctly specified 101 model. If the true residual plot exhibits noticeable deviations from these null plots, it 102 may suggest model misspecification. 103

Our computer vision model is trained to assign lower VSS to null plots and higher VSS to plots that display distinct patterns. Accordingly, statistical testing is performed by computing the proportion of null plots whose VSS equals or exceeds that of the

observed residual plot. This proportion serves as a p-value for a one-sided hypothesis test.

Bootstrapped residuals are obtained by refitting the model on bootstrap samples, 109 which are generated by sampling individual observations with replacement from the 110 original dataset. The residual plots obtained from these refitted models are evaluated 111 using the same computer vision model. The predicted VSS from the bootstrapped 112 plots provide an empirical estimate of the variation in the VSS of the observed 113 residual plot. By examining the proportion of bootstrapped plots that also exhibit 114 significant violations, we can assess whether the original conclusion is robust to 115 sampling variability. 116

# 4. R package: autovi

The main purpose of autovi is to provide rejection decisions and p-values for testing the null hypothesis  $(H_0)$  that the regression model is correctly specified. The package provides automated interpretation of residual plots using computer vision. The name autovi stands for automated visual inference. This functionality can be accessed through the R package autovi, or through a web interface, autovi.web, which enables users to perform analyses without installing R, Python, or their associated dependencies locally.

#### 4.1. Motivation

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Figure 1 shows three sets of plots of residuals against fitted values. The simulated 126 example in (a) might be interpreted as a heteroscedastic pattern, however the 127 automated reading would predict this to have a visual signal strength (VSS) of 128 1.53, with a corresponding p-value of 0.25. This means it would be interpreted as 129 a good residual plot, that there is nothing in the data to indicate a violation of 130 model assumptions. Skewness in the predictor variables is generating the apparent 131 heteroscedasticity, where the smaller variance in residuals at larger fitted values is 132 due to smaller sample size only. The Breusch-Pagan test (Breusch & Pagan 1979) for 133 heteroscedasticity would also not reject this as good residual plot. 134

The data in (b) is generated by fitting a linear model predicting mpg based on hp using
the datasets::mtcars. It is a small data set, and there is a hint of nonlinear structure
not captured by the model. The automated plot reading would predict a VSS of 3.57,
which has a p-value less than 0.05. That is, the nonlinear structure is most likely real,
and indicates a problem with the model. The conventional test, a Ramsey Regression

Equation Specification Error Test (RESET) (Ramsey 1969) would also strongly detect the nonlinearity.

The third example is generated using the surreal package (Balamuta 2024), where 142 structure residuals are embedded in the data. In this case, a quote inspired by Tukey, 143 'visual summaries focus on unexpected values', is used to define the residual structure. 144 The automated plot reading predicts the VSS to be 5.87, with a p-value less than 145 0.05. Visually, the structure is strikingly clear, but a RESET test for nonlinear 146 structure would not report a problem. (It would be detected by a Breusch-Pagan 147 for heteroscedasticity and also Shapiro-Wilk test (Shapiro & Wilk 1965) for non-148 149 normality.)

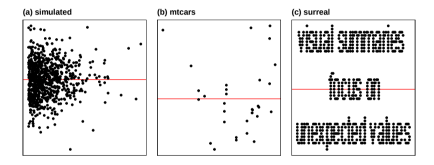


Figure 1. Reading residual plots can be a difficult task, particularly for students new to statistical modelling. The autovi package makes it easier. Here are three examples of residual plots, which may appear to have structure. According to autovi, the visual signal strengths (VSS) of these three examples are approximately (a) 1.53, (b) 3.57, (c) 5.87, resulting in (b), (c) being significant violations of good residuals, but (a) is consistent with a good residual plot.

#### 4.2. Implementation

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The autovi package is built on the bandicoot object-oriented programming (OOP) system (Li 2024), marking a departure from R's traditional S3 generic system. This OOP architecture enhances flexibility and modularity, allowing users to redefine key functions through method overriding.

The autovi infrastructure effectively integrates multiple programming languages and libraries into a comprehensive analytical tool. It relies on five core libraries from Python and R, each playing a critical role in the analysis pipeline. In Python, pillow (Clark et al. 2015) handles image processing tasks such as reading and resizing PNG files of residual plots, then converting them into input tensors for further analysis. TensorFlow

(Abadi et al. 2016), a key component of modern machine learning, is used to predict the VSS of these plots using a pre-trained convolutional neural network.

In the R environment, autovi utilizes several libraries. ggplot2 (Wickham 2016) generates the initial residual plots, saved as PNG files for visual input. cassowaryr (Mason et al. 2022) computes scagnostics (scatter plot diagnostics), providing numerical features that capture statistical properties of the plots. These scagnostics complement the visual analysis by offering quantitative metrics as secondary input to the computer vision model. reticulate (Ushey, Allaire & Tang 2024) enables seamless communication between R and Python.

#### 169 4.3. Installation

The autovi package is available on CRAN. It is actively developed and maintained, with the latest updates accessible on GitHub. This paper uses autovi version 0.4.2. The package includes internal functions to check the current Python environment used by the reticulate package. If the necessary Python packages are not installed in the Python interpreter, an error will be raised. If you want to select a specific Python environment, you can do so by calling the reticulate::use\_python() function before using the autovi package.

We recommend using the Shiny application autovi.web if users encounter installation problems.

# 179 4.4. Usage

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# 180 4.4.1. Numerical summary

Three steps are needed to get an automated assessment of a set of residuals and fitted values:

- 1. Load the autovi package using the library() function.
- 2. Create a checker object with a linear regression model.
- 3. Call the check() method of the checker, which, by default, predicts the VSS for the true residual plot, 100 null plots, and 100 bootstrapped plots. The method stores the predictions internally and prints a concise results report.
- 188 The code to do this is:

```
library(autovi)
checker <- residual_checker(lm(dist ~ speed, data = cars))
checker$check()</pre>
```

189 It produces the following summary:

```
<AUTO_VI object>
190
     Status:
191
      - Fitted model: lm
192
      - Keras model: (None, 32, 32, 3) + (None, 5) -> (None, 1)
193
         - Output node index: 1
194
      - Result:
195
         - Observed visual signal strength: 3.16 (p-value = 0.0396)
196
         - Null visual signal strength: [100 draws]
197
             - Mean: 1.274
198
             - Quantiles:
100
200
                   25%
                          50%
                                 75%
                                       80%
                                              90%
                                                     95%
                                                           99%
201
                 0.802 1.111 1.575 1.666 1.919 2.657 3.348
202
203
         - Bootstrapped visual signal strength: [100 draws]
204
             - Mean: 2.795 (p-value = 0.05941)
205
               Quantiles:
206
207
                   25%
                          50%
                                 75%
                                       80%
                                              90%
                                                     95%
                                                           99%
208
209
                 2.455 2.941 3.177 3.300 3.474 3.537 3.668
210
         - Likelihood ratio: 0.7333 (boot) / 0.06284 (null) = 11.67
211
```

212 The summary includes observed VSS of the true residual plot and associated p-value of the automated visual test. The p-value is the proportion of null plots (out of the 213 total 100) that have VSS greater than or equal to that of the true residual plot. The 214 report also provides sample quantiles of VSS for null samples and bootstrapped data 215 plots, providing more information about the sampling variability and a likelihood of 216 model violations. The likelihood is computed from the proportion of values greater 217 than the observed VSS in both the bootstrapped data values and the simulated null 218 values. 219

#### 4.4.2. Visual summary

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Users can visually inspect the original residual plot alongside a sample null plot using plot\_pair() or a lineup of null plot plot\_lineup(). This visual comparison can clarify why  $H_0$  is either rejected or not, and help identify potential remedies.

```
checker$plot pair()
```

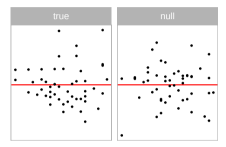


Figure 2. True plot alongside one null plot, for quick comparison.

The plot\_pair() method (Figure 2) displays the true residual plot on the left and a single null plot on the right. If a full lineup was shown, the true residual plot would be embedded in a page of null plots. Users should look for any distinct visual patterns in the true residual plot that are absent in the null plot. Running these functions multiple times can help any visual suspicions, as each execution generates new random null plots for comparison.

The package offers a straightforward visualisation of the assessment result through the summary\_plot() function.

```
checker$summary_plot()
```

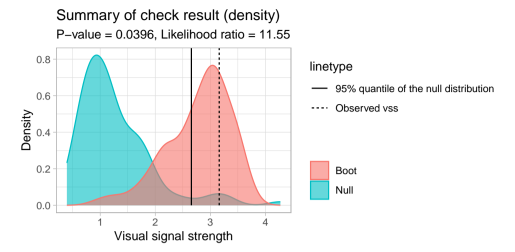


Figure 3. Summary plot comparing the densities of VSS for bootstrapped residual samples (red) relative to VSS for null plots (blue).

In the result, shown in Figure 3, the blue area represents the density of VSS for null residual plots, while the red area shows the density for bootstrapped residual plots. The dashed line indicates the VSS of the true residual plot, and the solid line marks the critical value at a 95% significance level. The p-value and the likelihood ratio are displayed in the subtitle. The likelihood ratio represents the ratio of the likelihood of observing the VSS of the true residual plot from the bootstrapped distribution compared to the null distribution.

Interpreting the plot involves several key aspects. If the dashed line falls to the right of the solid line, it suggests rejecting the null hypothesis. The degree of overlap between the red and blue areas indicates similarity between the true residual plot and null plots; greater overlap suggests more similarity. Lastly, the portion of the red area to the right of the solid line represents the percentage of bootstrapped models considered to have model violations.

This visual summary provides an intuitive way to assess the model's fit and potential violations, allowing users to quickly grasp the results of the automated analysis.

#### 4.5. Modularized infrastructure

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The initial motivation for developing autovi was to create a convenient interface for sharing the models described and trained in Li et al. (2024b). However, recognising that the classical normal linear regression model represents a restricted class of

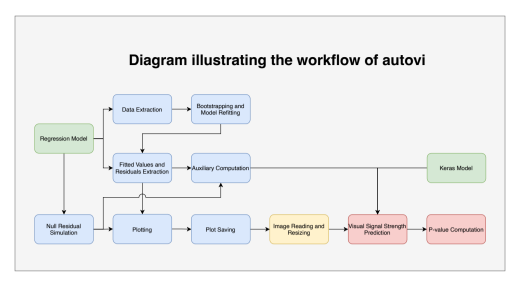


Figure 4. Diagram illustrating the infrastructure of the R package autovi. The modules in green are primary inputs provided by users. Modules in blue are overridable methods that can be modified to accommodate users' specific needs. The module in yellow is a pre-defined non-overridable method. The modules in red are primary outputs of the package.

models, we sought to avoid limiting the potential for future extensions, whether by
the original developers or other developers. As a result, the package was designed to
function seamlessly with linear regression models with minimal modification and few
required arguments, while also accommodating other classes of models through partial
infrastructure substitution. This modular and customisable design allows autovi to
handle a wide range of residual diagnostics tasks.

The infrastructure of autovi consists of ten core modules: data extraction, bootstrapping and model refitting, fitted values and residuals extraction, auxiliary computation, null residual simulation, plotting, plot saving, image reading and resizing, VSS prediction, and p-value computation. Each module is designed with minimal dependency on the preceding modules, allowing users to customise parts of the infrastructure without affecting its overall integrity. An overview of this infrastructure is illustrated in Figure 4.

The package takes regression models and a Keras model as primary inputs. Modules for VSS prediction and p-value computation are fixed but accessible via function arguments, using TensorFlow for inference and statistical testing. The image loading module is also fixed, using PIL to read and resize images based on the Keras model's input shape. The remaining seven modules are overridable, allowing users to adapt the workflow as needed. The data extraction module extracts a data.frame containing variables used in the regression model. The bootstrapping and refitting module resamples the data

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and refits the model. The fitted values and residuals extraction module returns these values as a data.frame. The auxiliary computation module calculates scagnostics such as monotonicity. The plotting module generates a ggplot in a standard format, and the plot saving module exports it at the same resolution as the training images.

These modules are described in detail in the package documentation.

#### 4.6. Extension to other model classes

- The autovi R package can be extended to accommodate other classes of models beyond linear regression, such as generalised linear models (glm). This is achieved by substituting the relevant overridable modules, and if needed, supplying a different Keras model.
- We provide an example of defining a new checker class tailored for Poisson regression using the glm framework:
  - 1. Define a new class using new\_class() with AUTO\_VI as the parent class.
    - 2. Override the necessary methods using register\_method(). In this example, we use Pearson residuals. To simulate null residuals, we assume the fitted model is correct and the estimated coefficients are accurate. New response values are generated accordingly, and a new model is fitted to this simulated response. Null residuals are then extracted from this refitted model.
    - 3. Create an alias for the instantiate() method of the new class.

The resulting checker class can be employed analogously to the linear model case described in Section 4.4. For illustration, we fit a Poisson model in which the quadratic term of the predictor x is intentionally omitted. This misspecification manifests as a pronounced U-shaped pattern in the lineap display (see Figure 5), which is also successfully identified by the computer vision model, yielding a p-value substantially below the conventional threshold of 0.05.

```
x <- rnorm(300, sd = 0.5)
y <- rpois(300, lambda = exp(1 + x + x^2))
pois_checker <- auto_pois_vi(
   glm(y ~ x, family = "poisson"),
   keras_model = get_keras_model("vss_phn_32")
)
pois_checker$plot_lineup()</pre>
```

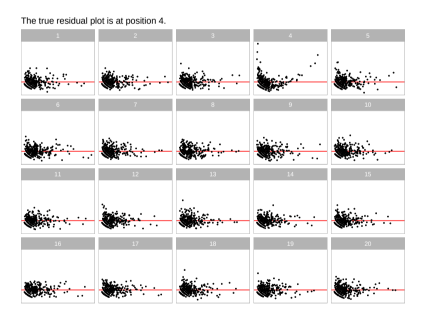


Figure 5. A linear of residual plots from Poisson generalised linear models, with the true residual plot at position 4, which displays a distinct U-shaped pattern. In contrast, the null plots show characteristics broadly consistent with well-behaved residuals from linear regression models.

#### pois\_checker\$check()

```
295 <AUTO_POIS_VI object>
296 Status:
```

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```
- Fitted model: glm, lm
297
      - Keras model: (None, 32, 32, 3) + (None, 5) -> (None, 1)
298
         - Output node index: 1
299
      - Result:
300
         - Observed visual signal strength: 4.875 (p-value = 0.009901)
301
         - Null visual signal strength: [100 draws]
302
             - Mean: 1.331
303
             - Quantiles:
304
305
                   25%
                          50%
                                 75%
                                       80%
                                              90%
                                                    95%
                                                           99%
306
                 1.035 1.233 1.488 1.644 1.941 2.276 2.639
307
308
         - Bootstrapped visual signal strength: [100 draws]
309
             - Mean: 5.51 (p-value = 0.009901)
310
             - Quantiles:
311
312
                          50%
                                 75%
                                       80%
                                              90%
                   25%
                                                    95%
                                                           99%
313
                 5.330 5.505 5.698 5.735 5.830 5.903 6.013
314
315
         - Likelihood ratio: 0.05096 (boot) / 0 (null) = Extremely large
316
```

It is important to note, however, that the pre-trained computer vision model included 317 in autovi, such as vss\_phn\_32 (see list\_keras\_models() for the full list of available 318 models), was developed specifically for diagnostics of linear regression. Its applicability 319 to other model classes relies on the assumption that the null residual plots exhibit 320 characteristics broadly consistent with those of well-behaved linear regression residuals, 321 that is, residuals should be approximately randomly scattered around zero, display 322 roughly constant variance across the range of fitted values, and exhibit no discernible 323 structure or curvature. If these conditions are not met, or if model violations do not 324 give rise to visually detectable patterns, the validity of the automated diagnostics may 325 be compromised. In such cases, users are encouraged to train and apply their own 326 Keras models. Detailed guidance on model training and discussion on extending the 327 methodology to other model classes can be found in Li et al. (2024b). 328

#### 5. Web interface: autovi.web

The autovi web Shiny application extends the functionality of autovi by offering a 330 user-friendly web interface for automated residual plot assessment. This eliminates the 331 common challenges associated with software installation, so users can avoid managing 332 Python environments or handling version requirements for R libraries. The platform 333 is cross-platform and accessible on various devices and operating systems, making it 334 suitable even for users without R programming experience. Additionally, updates are 335 managed centrally, ensuring that users always have access to the latest features. This 336 section discusses the implementation based on autovi.web version 0.1.0. 337

# 338 5.1. Implementation

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The interface autovi.web is built using the shiny (Chang et al. 2022) and shinydashboard (Chang & Borges Ribeiro 2021) R packages. Hosted on the shinyapps.io domain, the application is accessible through any modern web browser. The R packages htmltools (Cheng et al. 2024) and shinycssloaders (Sali & Attali 2020) are used to render markdown documentation in Shiny application, and for loading animations for Shiny widgets, respectively.

Determining the best way to implement the backend was difficult. In our initial planning for autovi.web, we considered implementing the entire web application using the webr framework (Moon 2020), which would have allowed the entire application to run directly in the user's browser. However, webr does not support packages which use compiled Fortran code, which is required by splancs (Rowlingson & Diggle 2023), a dependency of autovi. In the future, it is possible that a working Emscripten (Zakai 2011) version of this package may allow full webr support.

We also explored the possibility of implementing the web interface using frameworks 352 built on other languages, such as Python. However, server hosting domains that 353 natively support Python servers typically do not have the latest version of R installed. 354 355 Additionally, calling R from Python is typically done using the rpy2 Python library (Gautier 2024), but this approach can be awkward when dealing with language syntax 356 related to non-standard evaluation. Another option we considered was renting a server 357 where we could have full control, such as those provided by cloud platforms like Google 358 Cloud Platform (GCP) or Amazon Web Services (AWS). However, deploying and 359 maintaining the server securely requires some expertise. Ultimately, the most practical 360 solution was to use the shiny and shinydashboard frameworks, which are well-established 361 in the R community and offer a solid foundation for web application development. 362

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The server-side configuration of autovi.web is carefully designed to support its functionality. Most required Python libraries, including pillow and numpy, are preinstalled on the server. These libraries are integrated into the Shiny application using the reticulate package, which provides an interface between R and Python.

Due to shinyapps.io's resource policy, inactive servers enter sleep mode, clearing the local Python environment. When reactivated for a new session, libraries must be reinstalled. While this ensures a clean environment for each session, it may lead to slightly longer loading times for the first user after a period of inactivity.

In contrast to autovi, autovi.web leverages TensorFlow.js, a JavaScript library that allows
the execution of machine learning models directly in the browser. This choice enables
native browser execution, enhancing compatibility across different user environments,
and shifts the computational load from the server to the client-side. TensorFlow.js
also offers better scalability and performance, especially when dealing with resourceintensive computer vision models on the web.

While autovi requires downloading the pre-trained computer vision models from GitHub, these models in 'keras' file format are incompatible with TensorFlow.js. Therefore, we extract and store the model weights in JSON files and include them as extra resources in the Shiny application. When the application initialises, TensorFlow.js rebuilds the computer vision model using these pre-stored weights.

To allow communication between TensorFlow.js and other components of the Shiny application, the shinyjs R package (Attali 2021) is used. This package allows calling custom JavaScript code within the Shiny framework. The specialised JavaScript code for initialising TensorFlow.js and calling TensorFlow.js for VSS prediction is deployed alongside the Shiny application as additional resources.

#### 5.2. Usage

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The workflow of autovi.web is designed to be straightforward, with numbered steps displayed in each panel. There are two example datasets provided by the web application. The single residual plot example uses the dino dataset from the R package datasauRus (Davies, Locke & D'Agostino McGowan 2022). The lineup example uses residuals from a simulated regression model that has a non-linearity issue. We walk through the lineup example to further demonstrate the workflow of the web application.

#### 5.2.1. Reading data and setting parameters

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The user can select to upload data as either a single set of residuals and fitted values 395 in a two (or more) column CSV file or a pre-computed lineup of residuals and null 396 datasets in a three (or more) column CSV file (i.e. multiple sets of residuals and fitted 397 values with a column indicating the set label). Here we illustrate use with lineup 398 example data sets (Figure 6). To use the lineup example data, click the 'Use Lineup 399 Example' button. The data status will then update to show the number of rows and 400 columns in the dataset, and the CSV type will automatically be selected to the correct 401 option. Since the example dataset follows the variable naming conventions assumed 402 by the web application, the columns for fitted values, residuals, and labels of residual 403 plots are automatically mapped such that the column named as .fitted is mapped 404 to fitted values, .resid is mapped to residuals and if applicable, .sample to labels of 405 the residual set (middle image). If the user is working with a custom dataset, these 406 options must be set accordingly. Whenever a data containing a lineup, the user must 407 manually select the label for the true residual plot, otherwise the web application does 408 not provide all the results. The last step is to click the play button (right image) to **4**00 start the assessment. 410

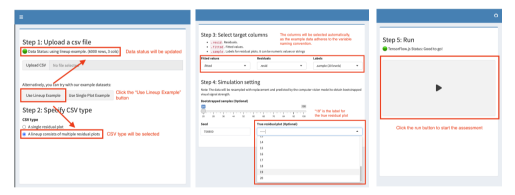


Figure 6. To begin the workflow for autovi using the lineup example dataset, the user clicks the 'Use Lineup Example' button (left) to load the example dataset, during which the data status and CSV type will be automatically updated. The user must manually select the label for the true residual plot (middle) to compute further results. The user initiates the assessment of the lineup example data by clicking the run button (right).

## 411 5.2.2. Results provided

Results are provided in multiple panels. The first row of the table Figure 7 is the most crucial to check, as it provides the VSS and the rank of the true residual plot among the other plots. The summary text beneath the table provides the *p*-value, which can be used for quick decision-making. The lineup is for manual inspection, and the user

should see if the true residual plot is visually distinguishable from the other plots, to confirm if the model violation is serious.

The density plot in Figure 8 offers a more robust result, allowing the user to compare the distribution of bootstrapped VSS with the distribution of null VSS. Finally, the grayscale attention map (right image) can be used to check if the target visual features, like the non-linearity present in the lineup example, are captured by the computer vision model, ensuring the quality of the assessment. The attention map is the gradient of the model output with respect to the grayscale image input, indicating the sensitivity of the output to each pixel.

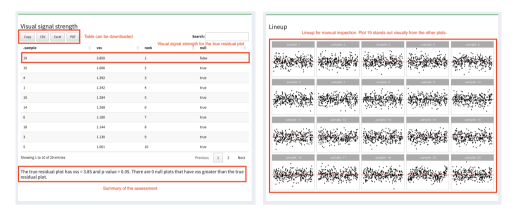


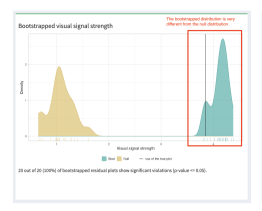
Figure 7. Results for the lineup. The VSS of the true residual plot is displayed in the first row of the table of VSS values for all the null plots (left image), with a summary text beneath the table providing the p-value to aid in decision-making. A lineup of residual plots allows for manual inspection (right image).

#### 6. Conclusions

This paper presents new regression diagnostics software, the R package autovi and its accompanying web interface, autovi.web. It addresses a critical gap in the current landscape of statistical software. While regression tools are widely available, effective and efficient diagnostic methods have lagged behind, particularly in the field of residual plot interpretation.

The autovi R package, introduced in this paper, automates the assessment of residual plots by incorporating a computer vision model, reducing reliance on time-consuming and potentially inconsistent human interpretation. This automation improves the efficiency of the diagnostic process and promotes consistency in model evaluation across different users and studies.

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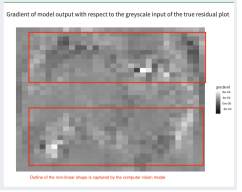


Figure 8. Summaries assessing the strength of the pattern and which elements of the plot contribute. The density plot helps verify if the bootstrapped distribution differs from the null distribution (left image). The attention map (right image) offers insights into whether the computer vision model has captured the intended visual features of the true residual plot.

The development of the accompanying Shiny app, autovi.web, expands access to these advanced diagnostic tools, by providing a user-friendly interface. It makes automated residual plot assessment accessible to a broader audience, including those who may not have extensive programming experience. This web-based solution effectively addresses the potential barriers to adoption, such as complex dependencies and installation requirements, that are often associated with advanced statistical software.

The combination of autovi and autovi.web offers a comprehensive solution to the challenges of residual plot interpretation in regression analysis. These tools have the potential to significantly improve the quality and consistency of model diagnostics across various fields, from academic research to industry applications. By automating a critical aspect of model evaluation, they allow researchers and analysts to focus more on interpreting results and refining models, rather than grappling with the intricacies of plot assessment.

The framework established by autovi and autovi.web opens up exciting possibilities for further research and development. Future work could explore the extension of these automated assessment techniques to other types of diagnostic plots and statistical models, potentially revolutionising how we approach statistical inference using visual displays more broadly.

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# 7. Resources and supplementary material

- The current version of autovi can be installed from CRAN, and source 455 code for both packages are available at github.com/TengMCing/autovi and 456 github.com/TengMCing/autovi web respectively. The web interface is available from 457 autoviweb.netlify.app. 458
- This paper is reproducibly written using Quarto (Allaire et al. 2024) powered by 450 Pandoc (MacFarlane, Krewinkel & Rosenthal 2024) and pdfTeX. The full source code 460
- to reproduce this paper is available at github.com/TengMCing/autovi\_paper. 461
- These R packages were used for the work: tidyverse (Wickham et al. 2019), Imtest 462
- (Zeileis & Hothorn 2002), kableExtra (Zhu 2021), patchwork (Pedersen 2022), rcartocolor 463
- (Nowosad 2018), glue (Hester & Bryan 2022), here (Müller 2020), magick (Ooms 2023), 464
- yardstick (Kuhn, Vaughan & Hvitfeldt 2024) and reticulate (Ushey, Allaire & Tang 465 2024).

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