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Felix Pretis†, Lea Schneider⋆, Jason E. Smerdon‡, and David F. Hendry†∗
†Department of Economics, and Institute for New Economic Thinking, Oxford Martin School, University of Oxford
⋆Department of Geography, Johannes Gutenberg University, Mainz
‡Lamont-Doherty Earth Observatory, Columbia University, Palisades, New York

Abstract

We present a methodology for detecting structural breaks at any point in time-series regression models using an indicator saturation approach. Building on recent developments in econometric model selection for more variables than observations, we saturate a regression model with a full set of designed break functions. By selecting over these break functions using an extended general-to-specific algorithm, we obtain unbiased estimates of the break date and magnitude. Monte Carlo simulations confirm the approximate properties of the approach. We assess the methodology by detecting volcanic eruptions in a time series of Northern Hemisphere mean temperature spanning roughly 1200 years, derived from a fully-coupled global climate model simulation. Our technique demonstrates that historic volcanic eruptions can be statistically detected without prior knowledge of their occurrence or magnitude- and hence may prove useful for estimating the past impact of volcanic events using proxy-reconstructions of hemispheric or global mean temperature, leading to an improved understanding of the effect of stratospheric aerosols on temperatures. The break detection procedure can be applied to evaluate policy impacts as well as act as a robust forecasting device.

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1 Overview

Breaks in time series come in many shapes and may occur at any point in time – distorting inference in-sample and leading to forecast failure out-of-sample if not appropriately modelled. Often an approximate shape of a break can be postulated *a priori*, either from previous observations or theory. For example, smooth transitions are common in economic time series following recessions or policy interventions, while sudden drops followed by smooth reversions to the mean are typical in climate time series such as temperature records after a large volcanic eruption (e.g., Kelly & Sear 1984). While the approximate form of a break may be known, the timings and magnitudes of breaks are often unknown. Here we propose an econometric approach for detecting breaks of any specified shape in regression models using an indicator saturation procedure. Our approach is based on recent developments in variable selection within regression models that involve more variables than observations (Castle et al. 2011). By selecting over a complete set of designed break indicators, our approach produces estimates of the break magnitude and timing without imposing limits on the number of breaks that may occur, even at the start or end of a sample.

A structural break is defined as a time-dependent change in a model parameter resulting from a change in the underlying data generating process. For example, a volcanic eruption leading to a rapid climatic cooling corresponds to a temporary shift in the mean of the surface temperature process. The detection of structural breaks in time series has received significant attention in the recent literature – with a growing interest in econometric models of climate change (e.g., Estrada et al. 2013, Pretis, Mann, & Kaufmann 2015). The focus has primarily remained on breaks in the mean through the form of step functions (Step-Indicator Saturation – SIS, Castle, Doornik, Hendry, & Pretis 2015, Pretis 2015b), smooth transition functions (González & Teräsvirta 2008), breaks in regression coefficients (see e.g. Bai & Perron 1998, 2003, Perron & Zhu 2005, Perron & Yabu 2009), or individual outliers or groups of outliers that can be indicative of different forms of breaks (Impulse-Indicator Saturation – IIS, see Hendry et al. 2008).

Broadly grouped into ‘specific-to-general’ and ‘general-to-specific’, there exist a plethora of approaches for the detection of structural breaks. Perron (2006) provides a broad overview of specific-to-general methods, some of which are subject to an upper limit on the number of breaks, a minimum break length, co-breaking restrictions, as well as ruling out breaks at the beginning or end of the sample, though Strikholm (2006) proposes a specific-to-general algorithm that allows for breaks at the start or end of a sample and relaxes the break-length assumption.

Indicator saturation (IIS, SIS) provides an alternative approach using an extended general-to-specific methodology based on model selection. By starting with a full set of step indicators in SIS and removing all but significant ones, structural breaks can be detected without having to specify a minimum break length, maximum break number or imposed co-breaking. Crucially this also allows model-selection to be conducted jointly with break-detection as non-linearities, dynamics, theory-motivated variables, and break functions are selected over simultaneously.

Step-functions and impulses are nevertheless only the simplest of many potential break specifications and may not provide the closest approximation to the underlying
break. Ericsson (2012) proposes a wide range of extensions to impulse and step-shifts. Here we show that the principle of step-indicator saturation can be generalized to any form of deterministic break function. An advantage over existing methods is an expected higher frequency of detection when a break function approximates the true break,\(^1\) high flexibility as multiple types of break functions can be selected over, and improvements in forecasting where designed functions act as continuous intercept corrections.

The method is illustrated using an econometric model of climate variation – detecting volcanic eruptions in a time series of Northern Hemisphere mean temperature spanning roughly 1200 years, derived from a fully-coupled global climate model simulation. Our technique demonstrates that eruptions can be statistically detected without prior knowledge of their occurrence or magnitude- and hence may prove useful for estimating the past impact of volcanic events using proxy-reconstructions of hemispheric or global mean temperatures. Specifically, this can lead to an improved understanding of the effect of stratospheric aerosols on temperatures (with relevance to geo-engineering and pollution control), and more generally, the break detection procedure can be applied to evaluate policy interventions (e.g., the Montreal Protocol: see Estrada et al. 2013, and Pretis & Allen 2013), correct measurement changes by detecting and subsequently removing shifts, and function as a robust forecasting device.

Section 2 introduces the methodology and investigates the properties of break detection in the presence of breaks and under the null of no breaks. Section 3 applies the method to detect volcanic eruptions in simulated climate data, and considers designed indicator functions as a robust forecasting device. Section 4 concludes.

## 2 Detecting breaks using designed indicator functions

Breaks are intrinsically stochastic without prior knowledge of their timings and magnitudes. Using a full set of break functions allows us to model the responses deterministically. The detection of structural breaks in regression models can be formulated as a model selection problem where we select over a full set of break functions, a subset of which accurately describes the underlying ‘true’ break. Consider a simple model as:

\[
y = Z\beta + \epsilon
\]

where \(y\) and \(\epsilon\) are \((T \times 1)\) vectors, \(\beta\) is a \((k \times 1)\) vector, and \(Z\) is a \((T \times k)\) matrix \(Z = (z_1, \ldots, z_k)\) of rank \(k\). We investigate the presence of structural breaks in any of the \(\beta\) where \(z\) may be a constant, trend or random variable. For each break type at any point in time for each variable whose coefficient is allowed to break, we augment the above model by a \((T \times T)\) break function matrix \(D\):\(^2\)

\[
y = Z\beta + D\gamma + \epsilon
\]

\(^1\)For example step-indicator saturation exhibits higher power in detecting step shifts than using impulses alone – see Castle, Doornik, Hendry, & Pretis (2015).

\(^2\)For \(k\) breaking variables this implies augmenting the full-sample model by \(k\) \((T \times T)\) matrices.
where $\gamma$ is a $(T \times 1)$ vector. The specification of $D$ is such that the first column $d_1$ $(T \times 1)$ is set to denote some specified break function $d(t)$ of length $L$, where $d_{1,t} = d(t)$ for $t \leq L$ and 0 otherwise, $d_{1,t} = 0$ for $t > L$. All further columns $d_j$ (for $j = 2, ..., T$) in $D$ are set such that $d_{j,t} = d_{j-1,t-1}$ for $t \geq j$ and 0 otherwise. The break matrix $D$ is then defined as $D = (d_1, d_2, ..., d_T)$, where $d_j$ denotes a vector with break at time $t = j$:

$$D = (d_1, d_2, ..., d_T)$$

$$d_1 = (d_1, d_2, ..., d_{L-1}, d_L, 0, ..., 0)'$$

$$d_2 = (0, d_1, d_2, ..., d_{L-1}, d_L, 0, ..., 0)'$$

$$d_3 = (0, 0, d_1, d_2, ..., d_{L-1}, d_L, 0, ..., 0)'$$

$$...$$

This specification provides a general framework within which multiple break types can be analysed – Table 1 provides a non-exhaustive overview. The form of the break function $d(t)$ has to be designed a-priori, but this is implicitly done in most structural break detection methods. For example, outlier detection through finding impulses (IIS, in Hendry et al. 2008) sets the break vector in $d_1$ such that $d(t) = 1$ and $L = 1$, while a search for step-shifts (SIS) sets $d(t) = 1$ and the length to $T - t + 1$, i.e. the break function continues until the end of the sample. Breaks in linear trends (see e.g., Perron & Yabu 2009, Perron & Zhu 2005, Estrada et al. 2013) can be constructed by setting $d(t) = t$ and the length to $T - t + 1$. Pretis, Mann, & Kaufmann (2015) apply indicator saturation using broken trends and step shifts to evaluate climate models. Breaks in coefficients on random variables $z_t$ (see e.g. Bai & Perron 2003, Ericsson 2012, Kitov & Tabor 2015) can be constructed by interacting $z_t$ with a full set of step shifts. Sudden

$3$While the framework presented here provides an encompassing specification for many break types, the construction of $D$ is not limited to this particular case. Additional sets of specifications for step shifts are considered in Castle, Doornik, Hendry, & Pretis (2015). The appeal of the specification here is that the definition of $D$ allows for a general framework under which properties can be analyzed where many of the previously proposed cases are a special case of $D$. 

<table>
<thead>
<tr>
<th>Break value: $d(t)$</th>
<th>Length:</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Deterministic Breaks</strong></td>
<td></td>
</tr>
<tr>
<td>General Case</td>
<td>$d(t)$</td>
</tr>
<tr>
<td>Impulses (IIS)</td>
<td>1</td>
</tr>
<tr>
<td>Step Shifts (SIS)</td>
<td>1</td>
</tr>
<tr>
<td>Broken Trends</td>
<td>$t$</td>
</tr>
<tr>
<td>Volcanic Functions</td>
<td>see eq. (31)</td>
</tr>
<tr>
<td><strong>Random Variables</strong></td>
<td></td>
</tr>
<tr>
<td>Coeff. on $z_t$ (MIS)</td>
<td>$z_t \cdot d_{t,SIS}$</td>
</tr>
</tbody>
</table>

4
declines followed by a smooth recovery to the mean in hemispheric temperature responses are introduced here as volcanic functions and considered in sections 2.1 and 3. Linear combinations of multiple break functions can allow for varying lengths of breaks without pre-specification.

Searching for breaks in \( k \) variables implies that the complete break matrix across all \( k \) variables \( D \) is of dimension \( (T \times kT) \). The inclusion of \( kT \) additional variables leads to the total number of variables \( N \) exceeding the number of observations, \( N > T \), even for \( k = 1 \). Thus, a methodology allowing for more variables than observations is required.

Selection of models with more variables than observations has primarily relied on either shrinkage-based penalised likelihood methods (Tibshirani 1996, Zou & Hastie 2005, Tibshirani 2011) or general-to-specific methodology in the econometrics literature (see e.g. Castle et al. 2011). Kock & Teräsvirta (2015) compare the general-to-specific model selection algorithms Autometrics, to QuickNet – an artificial neural-network method proposed by H. White (2006), and to a shrinkage-based bridge estimator from Huang et al. (2008) in the context of forecasting.

Here we rely on general-to-specific model selection due to methods based on forward-step-wise searches not performing as well in break detection contexts (see section 2.1.2 for a simple comparison, or Epprecht et al. n.d., and Hendry & Doornik 2014 for comparisons on general variable selection.). Cox and Snell (1974) discuss some of the challenges of the general variable selection problem and Hoover & Perez (1999) show the feasibility of general-to-specific model selection for \( N << T \). When facing more variables than observations, the general-to-specific approach is closely linked to robust statistics. Saturating a model with a full set of \( 0/1 \) indicator functions from which selections are made is equivalent to a robust one-step M-estimator using Huber’s skip function (see Johansen & Nielsen 2009, Johansen & Nielsen 2013 for the iterated case, and Johansen & Nielsen 2016 for an overview). Here we generalize this allowing for any form of designed break function in place of impulses, and formulate break detection as a model selection problem.

To estimate model (2) saturated with a full set of break functions \( D \) (so \( N > T \)), we rely on a block-partitioning estimation procedure (Doornik 2010, Hendry & Johansen 2015). For this, we partition \( D \) into \( b \) blocks of \( n_i \) variables such that \( n_i << T \) and \( \sum_{i=1}^{b} n_i = N \). In the simplest case of testing for a break in a single variable (e.g. the intercept), a split-half approach (see Figure 1 and Algorithm 1 in supplementary material 5.4) is feasible: initially we include the first half of \( D_1 = (d_1, \ldots, d_{T/2}) \) and retain only significant break indicators. We repeat the step for the second half of break functions \( d_j \) (for \( j = T/2 + 1, \ldots, T \)) and finally combine the retained sets and only keep significant indicators. This split-half approach is considered here for analytical tractability in section 2.1.2.

In practice, however, we rely on a multi-split and multi-path search to lower the variance of the estimators, allow for any number of variables for a given set of observations, and to avoid a breakdown of the procedure if the breaks cannot be adequately modelled through split-half indicators.\(^4\) This can be implemented through the

\(^4\)In a simple split-half analysis there may be an identification problem if the sample-split coincides
general-to-specific model selection algorithm *Autometrics* (Algorithm 2 in supplementary material 5.4), described in Doornik 2009a, or the *gets* package in the statistical software environment *R* (Pretis, Sucarrat, & Reade 2015). The algorithm (referred to as multi-path throughout the paper) avoids path dependence through a tree-structure and uses a parallel step-wise backwards search, while testing for encompassing and congruence (Hendry 1995). See Hendry & Pretis 2013 for an application of the algorithm to econometric modelling of climate change. A simulation-based comparison to shrinkage methods is provided in section 2.1.2.

2.1 Properties of designed indicator functions in the presence of breaks

To assess the theoretical power of the proposed methodology, we first investigate the properties in the benchmark case of a single break matched by a correctly timed break indicator in section 2.1.1. Section 2.1.2 then assesses the properties of break indicator saturation when the break date and magnitude are unknown. Section 2.1.3 investigates uncertainty around the break date and 2.2 describes the properties in the presence of no breaks. Theory results are derived for general designed functions, simulation examples are based on a volcanic break as characterised by equation (31) below.

2.1.1 Power for a known break date

We investigate the theoretical power of detecting a break in a time series given a known break date. Consider a data generating process (DGP) coinciding with the model for a single known break in an intercept:

\[ y_t = \mu + \lambda d_t + \epsilon_t \]  

(4)

where \( \epsilon_t \sim \text{IN}(0, \sigma^2) \). The break shifts \( \mu \) to \( \mu + \lambda d_t \) where \( d_t \) is a break function of length \( L \) beginning at time \( t = T_1 \) where \( (T_1 + L) \leq T \) such that \( d_t \neq 0 \) for \( T_1 \leq t < (T_1 + L) \) and 0 otherwise. The estimators \( \hat{\mu} \) and \( \hat{\gamma} \) (where \( \hat{\gamma} \) is the estimator for \( \lambda \)) in a correctly specified model for a known break are given by:

\[
\begin{bmatrix}
\hat{\mu} - \mu \\
\hat{\gamma} - \lambda
\end{bmatrix} =
T_d^{-1} \begin{bmatrix}
\sum_{t=T_1}^{T_1+L-1} d_t^2 \sum_{t=1}^{T} \epsilon_t - \sum_{t=T_1}^{T_1+L-1} d_t \sum_{t=T_1}^{T} \epsilon_t \\
\sum_{t=T_1}^{T_1+L-1} d_t \sum_{t=1}^{T} \epsilon_t - \sum_{t=T_1}^{T_1+L-1} d_t \sum_{t=0}^{T} \epsilon_t
\end{bmatrix}
\]  

(5)

where \( T_d = T \left[ \sum_{t=T_1}^{T_1+L-1} d_t^2 - \frac{1}{T} \left( \sum_{t=T_1}^{T_1+L-1} d_t \right)^2 \right] \). The estimators are unbiased for the break and intercept: \( E[\hat{\mu} - \mu] = 0 \) and \( E[\hat{\gamma} - \lambda] = 0 \). The variance of the estimators is given by:

\[
V \begin{bmatrix}
\hat{\mu} - \mu \\
\hat{\gamma} - \lambda
\end{bmatrix} = \sigma^2 T_d^{-1} \begin{bmatrix}
\sum_{t=T_1}^{T_1+L-1} d_t^2 - \sum_{t=T_1}^{T_1+L-1} d_t T \\
- \sum_{t=T_1}^{T_1+L-1} d_t T
\end{bmatrix}
\]  

(6)

perfectly with a structural break. This is overcome by varying the block partitioning as is done in the software implementations of the algorithm.
The distribution of the break estimator is then:

\[
(\hat{\gamma} - \lambda) \sim N \left(0, \sigma^2 \left[ \sum_{t=T_1}^{T_1+L-1} d_t^2 - \sum_{t=T_1}^{T_1+L-1} d_t \bar{d} \right]^{-1} \right)
\]  

(7)

where \( \bar{d} = 1/T \sum_{t=1}^{T} d_t \). For the special case when step-indicators are chosen as the functional form of \( d_t \) and the single break lasts from \( t = 0 \) to \( t = T_1 < T \), equation (5) simplifies to \( \hat{\mu} - \mu = \bar{\epsilon}_2 \) and \( \hat{\gamma} - \lambda = \bar{\epsilon}_1 - \bar{\epsilon}_2 \), where \( \bar{\epsilon}_1 = 1/T_1 \sum_{t=1}^{T_1} \epsilon_t \) and \( \bar{\epsilon}_2 = 1/(T - T_1) \sum_{t=T_1+1}^{T} \epsilon_t \).

### 2.1.2 Potency for an unknown break date

When the break date is unknown, we propose to saturate the regression model using a full set of specified break indicators and select significant breaks through an extended general-to-specific algorithm (Castle et al. 2011). We assess the methodology in the selection context using two main concepts: the null retention frequency of indicators is called the gauge, comparable to the size of a test denoting its (false) null rejection frequency, but taking into account that indicators that are insignificant on a pre-assigned criterion may nevertheless be retained to offset what would otherwise be a significant misspecification test (see Johansen & Nielsen 2016 for distributional results on the gauge). The non-null retention frequency when selecting indicators is called its potency, comparable to a similar tests power for rejecting a false null hypothesis.

Here we investigate the feasibility of the proposed method by deriving the analytical properties of the split-half approach for an unknown break. Figure 1 illustrates the split-half method for a single unknown break. In practice we rely on a multi-path, multi-block search algorithm (such as Autometrics, see Algorithm 2 in section 5.4) to reduce the variance of the estimators.

Consider a single break falling into the first half of the sample beginning at time \( T_1 \) for \( L \) periods such that \( 0 < T_1 < T_1 + L < T/2 \). In matrix form the DGP is given as:

\[
y = \lambda d_{T_1} + \epsilon
\]  

(8)

where \( \epsilon \sim \text{IN}(0, \sigma^2) \) for simplicity and the \((T \times 1)\) vector \( d_{T_1} \) denotes a break at \( t = T_1 \) for \( L \) periods. Using a split-half approach, we assess the properties of detecting the single break when the break date is unknown. The split-half model for the first half of break functions is:

\[
y = D_1 \gamma_{(1)} + \nu
\]  

(9)

where \( \gamma_{(1)} = (\gamma_1, \gamma_2, ... \gamma_{T/2})' \) and \( D_1 = (d_1, ..., d_{T/2}) \). The estimator \( \hat{\gamma}_{(1)} \) equals:

\[
\hat{\gamma}_{(1)} = (D_1' D_1)^{-1} D_1' y = \lambda (D_1' D_1)^{-1} D_1' d_{T_1} + (D_1' D_1)^{-1} D_1' \epsilon
\]  

(10)

5See supplementary material 5.3 for proof.

6Proof given in supplementary material 5.2.
Figure 1: Split-half approach for a single unknown break of the shape of a volcanic function at T=75. Left column shows included indicators in each step, middle column shows the retained indicators, right column graphs the selected model with actual and fitted data. Block 1 (top panel) includes the first half of break functions and retains a single one as the mean is lowered in the second half due to the presence of a break at t=75. Block 2 (middle panel) then includes the second half retaining the correct break function. Block 3 uses the union of retained indicators from blocks 1 and 2 in which now the first indicator is rendered insignificant by the mean being correctly estimated due to the second indicator capturing the break. Using a saturating set of break functions at 1% the break at T=75 is detected without prior knowledge and is the only break function retained.
where the $(T/2 \times 1)$ vector $r$ is equal to one at $t = T_1$ and zero otherwise, $r_t = 1_{\{t=T_1\}}$. It follows that $E[\gamma(1)] = \lambda r$ and $V[\gamma(1)] = \sigma_r^2 (D'_1 D_1)^{-1}$. We find for the first half, for normal error terms:

$$
(\hat{\gamma}_1 - \lambda r) \sim N \left( 0, \sigma_r^2 (D'_1 D_1)^{-1} \right) \quad (11)
$$

Therefore conventional t-tests can be used to assess the significance of individual indicators. The estimator $\hat{\gamma}(2)$ on the second half of indicators, $D_2 = (d_{T/2+1}, \ldots, d_T)$, will miss the break in the DGP in the first half described by $d_{T_1}$ and equals:

$$
\hat{\gamma}(2) = \lambda (D'_2 D_2)^{-1} D'_2 d_{T_1} + (D'_2 D_2)^{-1} D'_2 \epsilon \quad (12)
$$

For step-shifts, Castle, Doornik, Hendry, & Pretis (2015) show that the indicator in $D_2$ closest to the sample split will be retained in the second set of indicators. For the general form of break functions, retention in $D_2$, when there is a break in the first half, will depend on the specific functional form. However, conditional on the break indicator being correctly retained in the first set $D_1$, retention of irrelevant indicators in $D_2$ does not affect the correct identification of the break overall: let $D_{1*}$ and $D_{2*}$ denote the set of retained break functions in the first and second set respectively, where retention is based on a retention rule such as $d_j$ is retained if $|t_{\hat{\gamma}_j}| \geq c_\alpha$. The final step in the split-half procedure is then to combine the retained indicators using $D_U = [D_{1*} D_{2*}]$ and estimate the model:

$$
y = D_U \gamma(U) + \nu \quad (13)
$$

This yields the estimator $\hat{\gamma}(U)$ unbiased for the true break:

$$
\hat{\gamma}(U) = \lambda r + (D'_U D_U)^{-1} D'_U \epsilon \quad (14)
$$

The carried-forward break function in $D_{1*}$ correctly identifies the true break, and coefficients on all other break functions will thus be zero in expectation. The proof is identical to that given for the first half of indicators in the supplementary material 5.2. This shows that, conditional on retaining the correct break indicator in $D_1$, the retention of indicators in $D_2$ does not affect the correct identification of the break, when the first and second set are combined and re-selected over. The distribution of the final split-half estimator is then given by:

$$
(\hat{\gamma}(U) - \lambda r) \sim N \left( 0, \sigma^2_s (D'_U D_U)^{-1} \right) \quad (15)
$$

Re-selection then results in only the true break indicator being retained in expectation.\footnote{In practice selection bias can be controlled using bias-correction after orthogonalization of the selected regressors – see Hendry & Krolzig (2005) for the orthogonal case, Pretis (2015b) for bias-correction of step-functions, and Castle, Doornik, & Hendry (2015) for bias-correction with correlated variables.}

This result generalizes the specific case of step indicators presented in Castle, Doornik, Hendry, & Pretis (2015). Even though the break date and magnitude are unknown, the
use of a fully saturated set of break indicators allows us to obtain an unbiased estimate of the break magnitude and timing. The estimator then follows a normal distribution subject to correct specification of the break function. Thus the estimated coefficient at the break time, $\hat{\gamma}_{T_1}$, is in expectation equal to the break magnitude, while all other estimated coefficients are mean-zero in expectation. This result generalizes to multiple breaks falling in a single split. As in the case of the known break timing, the variance of the estimator depends on the specified break function. Let $\delta_{k,j}$ denote the $(k,j)$ element of the matrix $(D_1' D_1)^{-1}$. The variance of the coefficient at the breakpoint in the first half is therefore:

$$V[\hat{\gamma}_{T_1}] = \sigma^2 \delta_{T_1,T_1}$$  \hspace{1cm} (16)

For iid error terms $\epsilon$, and $D$ specified as a full set of step functions, the split-half model (without selection) yields $\delta_{j,j} = 2$, so the break coefficient has twice the error variance. For the proposed volcanic function (derived and assessed in detail in section 3) modelling a single drop followed by a reversion to the mean, we find that $\delta_{j,j} = 3.7$, thus $V[\hat{\gamma}_{T_1}] = 3.7\sigma^2$. This can be compared to the known-break/single-indicator case where the variance is given by equation (6) and for the volcanic function equals $2.3\sigma^2$ (for $T=100$). Due to collinearity of break functions, the variance of the estimator is higher in a fully saturated model. In the more general case, $\delta_{T_1,T_1}$ depends on the specification of the break function but can be computed a-priori. The $t$-statistic is then given as:

$$t_{\hat{\gamma}_{T_1}} = \frac{\hat{\gamma}_{T_1}}{\epsilon \sqrt{\delta_{T_1,T_1}}} \approx \frac{(\hat{\gamma}_{T_1} - \lambda)}{\epsilon \sqrt{\delta_{T_1,T_1}}} + \frac{\lambda}{\epsilon \sqrt{\delta_{T_1,T_1}}} \sim N \left( \frac{\lambda}{\epsilon \sqrt{\delta_{T_1,T_1}}}, 1 \right)$$  \hspace{1cm} (17)

In practice we use sequential elimination of the break indicators or a multi-path search to eliminate insignificant indicators reducing the variance of the estimators from a saturated model (16) closer to the single break (6) and increasing the power of detection.

For dynamic time series models, the above approach can be extended by including time-dependent covariates. Valid conditioning (e.g. through the inclusion of autoregressive terms in the case of non-iid errors) can be ensured by always including the covariates in each block estimation step and only selecting over the break functions. Johansen & Nielsen (2009) provide the asymptotics under the null of no break for the special case of impulses for stationary and unit-root non-stationary autoregressive processes (see Johansen & Nielsen 2013, for the iterated version). The case for general break functions is discussed in section 2.2, and the supplementary material 5.5 provides simulation results for an AR(1) model and DGP.

9While our analysis concentrates on small-sample properties, the asymptotic rates of convergence will generally depend on the specification of the break function – varying scaling to obtain non-degenerate limit distributions may therefore be required. In the case of step functions ($d_t = 1, L = T$) and the simple no-intercept case, pre-multiplying the estimator by $\sqrt{T}$ yields asymptotic normality for the break estimator when $T^{-1} \sum_{t=1}^{T-1} d_t^2 = T^{-1} L \to \tau$ as $T \to \infty$. In other words, the ratio of break length to the sample size remains constant as the sample size increases – this can be interpreted as obtaining more information on the break period or sampling at higher frequencies as $T \to \infty$. A similar analysis can be applied to the volcanic functions considered here, where either the break length scales with the sample size, or alternatively the magnitude increases similar to the asymptotic analysis for a single impulse in Doornik et al. (1998).
Simulation performance based on volcanic break functions: Table 2 reports simulation results (T=100) for a DGP with a single unknown volcanic break at $t = T_1 = 25$ of magnitude $\lambda$ followed by a smooth reversion to the mean.\textsuperscript{10} Equation (31) provides the exact functional form. Simulations are assessed by the retention/detection frequency (potency) for a single break and average retention of spurious breaks (gauge).\textsuperscript{11}

The trade-off between potency and level of significance of selection $\alpha$ is shown in Figure 2 for a single volcanic break. A multi-path search generally increases the power of detection relative to the split-half approach. Figure 3 shows the results for split-half (dashed) and multi-path (solid) selection when using volcanic functions for a break of $\lambda = 6$. Consistent with derived theory (16), the estimator has 3.7 times the variance of the error term when using split-half estimation for the given function. Using a multi-path search reduces the variance drastically. Any selection bias of the multi-path search estimates can be controlled through bias correction after selection (see Castle et al. 2011 and Pretis 2015b). The supplementary material 5.5 provides simulation results for a simple autoregressive DGP and model.

**Table 2:** Potency of detecting an unknown break when using split-half and multi-path searches. Statistics were generated from 1000 simulations and detection significance was set to $\alpha = 0.01$, with a length of $L = 3$. Break magnitude $\lambda$ corresponds to the full response in standard deviations of the error term ($\sigma = 1$) over the entire break, the trough is $0.58\lambda$.

<table>
<thead>
<tr>
<th></th>
<th>Split-Half</th>
<th>Multi-Path</th>
</tr>
</thead>
<tbody>
<tr>
<td>$\lambda = 6$, trough=3.48</td>
<td>Potency 0.69, Gauge D1 0.013</td>
<td>Potency 0.88, Gauge 0.015</td>
</tr>
<tr>
<td>$\lambda = 4$, trough=2.23</td>
<td>Potency 0.30, Gauge D1 0.013</td>
<td>Potency 0.50, Gauge 0.014</td>
</tr>
<tr>
<td>$\lambda = 2$, trough=1.16</td>
<td>Potency 0.06, Gauge D1 0.013</td>
<td>Potency 0.11, Gauge 0.015</td>
</tr>
</tbody>
</table>

**Comparison to shrinkage-based methods** Shrinkage-based methods using penalised likelihood estimation (Tibshirani 2011, Zou & Hastie 2005) provide an alternative to the general-to-specific algorithm used here in selecting models with more variables than observations. Figure 4 shows the simulation outcomes comparing multi-path indicator saturation (for $\alpha = 0.01$), the Lasso (Tibshirani 1996, estimated using LARS, see Efron et al. 2004) where cross-validation is used to determine the penalty, and the Lasso where the penalty is set such to approximate the false-positive rate of the IS procedure under the null of no breaks ($\approx 0.01$). The simulation uses a total break magnitude of 6 standard deviations (implying a trough of $3.48\sigma$) for an increasing number of evenly spaced breaks from 0 up to 10 in a sample of $T = 100$. The general-to-specific multi-path algorithm exhibits stable power exceeding that of the penalised likelihood

\textsuperscript{10}For a volcanic break, $\lambda$ denotes the entire temperature response over the specified length $L$, thus the trough will be less than $\lambda$. For the present specification of $L = 3$, the initial trough of the function equals $0.58\lambda$.

\textsuperscript{11}All simulations and applications using the multi-path search Autometrics are coded using the Ox programming language (Doornik 2009b). Simulations using the Lasso are coded using the package glmnet (Friedman et al. 2010) in R.
Figure 2: Left: Potency of detecting a volcanic break of magnitude \( \lambda \) for level of significance \( \alpha \) using split-half and multi-path selection. Right: Proportion of spuriously retained break indicators (gauge). Break magnitude \( \lambda \) corresponds to the full response in standard deviations of the error term \( \sigma_e = 1 \) over the entire break, the peak is \( 0.58\lambda \).

Figure 3: Estimated break indicator and variance for unknown break using split-half (blue) and multi-path (yellow) selection. The left panel shows a simulated time series with the true break shown in blue and the fit in green. The middle panel shows the distribution of the estimated coefficient, the right panel shows the variance of the coefficient. Dashed lines show the true break magnitude and analytical variance of the split-half coefficient.
Figure 4: Left: Average potency of detecting increasing numbers of volcanic breaks using multi-path indicator saturation at $\alpha = 0.01$ (IS, orange), cross-validated Lasso (blue), and Lasso with fixed penalisation parameter (green) where the penalisation parameter is set such that the false-positive rate approximates that of the indicator saturation procedure under the null of no break. Right: Corresponding false-positive rate (gauge). $M = 1000$ replications.

methods across any number of breaks. The false-positive rate remains stable and close to the theory level of 0.01. The shrinkage-based procedures, due to their similarity to forward-selection, show decreasing potency as the number of breaks increases, and the false-positive rate is difficult to control.

2.1.3 Uncertainty around the break date

An estimated uncertainty on the break magnitude and coefficient path (the time-varying intercept in the regression) can be computed given the distribution of the break estimator (see Pretis 2015b). While of considerable interest, it is non-trivial, however, to quantify the uncertainty around the timing of the break (see Elliott & Müller 2007). This is particularly true for the literature focusing on break detection using general to specific methodology. Here we investigate the uncertainty around the timing of estimated break points when using break indicator saturation by computing the analytical power of a single break indicator when the break function is correctly specified but the break time is not. This is a simplification as it only considers a single mis-timed indicator, while the indicator saturation approach includes a saturating set.

Consider a DGP with just a single break in the mean:

$$y_t = \lambda d_{T_1,t} + \epsilon_t$$  \hspace{1cm} (18)

The break shifts $E[y_t]$ from 0 to $\lambda d_{T_1}$ at $t = T_1$ where $d_{T_1}$ is a break function of length $L$ beginning at time $t = T_1$ such that $T_1 + L < T$ and $d_{T_1} = (0, ..., d_1, d_2, ..., d_L, 0, ..., 0)$. The corresponding model is then:

$$y_t = \gamma d_{j,t} + \nu_t$$  \hspace{1cm} (19)
When the break date is correctly specified \( d_{j,t} = d_{T_1,t} \), so the estimator for \( \lambda \) is given by:

\[
\hat{\gamma}_{t=T_1} - \lambda = \left( \sum_{t=T_1}^{T_1+L} d_{T_1,t}^2 \right)^{-1} \left( \sum_{t=T_1}^{T_1+L} d_{T_1,t} \right) \tag{20}
\]

Similarly for a test of the hypothesis: \( \lambda = 0 \), the \( t \)-statistic has a noncentrality of

\[
E [t_{\hat{\gamma},t=T_1}] = \psi = \frac{\lambda \sqrt{\left( \sum_{t=T_1}^{T_1+L} d_{T_1,t}^2 \right)}}{\sigma_e} \quad \text{and the normal distribution:}
\]

\[
t_{\hat{\gamma}} \approx \frac{\hat{\gamma}_{t=T_1} \sqrt{\left( \sum_{t=T_1}^{T_1+L} d_{T_1,t}^2 \right)}}{\sigma_e} \sim N(\psi,1) \tag{21}
\]

The noncentrality \( \psi \) increases in the break magnitude \( \lambda \), varies with the break length \( L \), and will depend on the underlying break function given by \( d_t \).

Now consider the model being incorrectly specified for the break date, such that \( d_{j,t} \neq d_{T_1,t} \) but is shifted by \( K \) periods \( d_{j,t} = d_{T_1 \pm K,t} \). The estimator for \( \lambda \) is then:

\[
\hat{\gamma}_{t=T_1 \pm K} - \lambda = \lambda \left[ \left( \sum_{t=T_1}^{T_1+L} d_{j,t}^2 \right)^{-1} \left( \sum_{t=T_1}^{T_1+L} d_{j,t}d_{T_1,t} \right) - 1 \right] + \left( \sum_{t=T_1}^{T_1+L} d_{j,t} \right) \left( \sum_{t=T_1}^{T_1+L} d_{j,t} \right)^{-1} \left( \sum_{t=T_1}^{T_1+L} d_{j,t} \right) \tag{22}
\]

For a fixed length \( L \) and a forced mis-timing, it follows that \( \hat{\gamma}_{t \neq T_1} \) is not an unbiased estimator for \( \lambda \). Note that if \( d_j \) is functionally specified correctly such that the only difference to the true break function is through \( K \) lags, \( d_j = d_{T_1 \pm K} \), then it holds that

\[
\left( \sum_{t=T_1}^{T_1+L} d_{j,t}^2 \right) = \left( \sum_{t=T_1}^{T_1+L} d_{T_1,t}^2 \right) \quad \text{Equally } \left( \sum_{t=T_1}^{T_1+L} d_{j,t}d_{T_1,t} \right) = \left( \sum_{t=T_1}^{T_1+L} d_{T_1 \pm K,t}d_{T_1,t} \right)
\]

for \( K \leq L \) and 0 for \( K > L \). Using this, we derive an expression for the approximate \( t \)-statistic associated with the estimator given a break function time mis-specified by \( K \) lags:

\[
E [t_{\hat{\gamma},t=T_1 \pm K}] \approx \frac{E [\hat{\gamma}_{t=T_1 \pm K}]}{\sigma_e \left( \sum_{t=T_1}^{T_1+L} d_{T_1,t}^2 \right)^{-1/2} } = \frac{\lambda \left( \sum_{t=T_1}^{T_1+L} d_{T_1 \pm K,t}d_{T_1,t} \right)}{\sigma_e \left( \sum_{t=T_1}^{T_1+L} d_{T_1,t}^2 \right)^{1/2} } \tag{23}
\]

This is equal to the noncentrality of the correct break date \( \psi \) scaled by a factor less than one, decreasing with the distance \( K \) from the correct date:

\[
E [t_{\hat{\gamma},t=T_1 \pm K}] \approx \psi \left( \frac{\sum_{t=T_1}^{T_1+L} d_{T_1 \pm K,t}d_{T_1,t}}{\sum_{t=T_1}^{T_1+L} d_{T_1,t}^2} \right) \leq \psi \tag{24}
\]

For a given break specification \( d_t \) and break length \( L \), the corresponding power function can be computed to provide an approximate measure of power for detection of a break at \( t = T_1 \) in the neighbourhood of \( T_1 \). Note that \( E [t_{\hat{\gamma},t=T_1 \pm K}] \) is zero outside a neighbourhood of \( L \). The associated \( t \)-statistic of a break indicator further away from the true break date \( T_1 \) than the break length \( L \) is zero in expectation, since

\[
\left( \sum_{t=T_1}^{T_1+L} d_{j,t}d_{T_1,t} \right) = 0 \quad \text{for } K > L
\]

Intuitively, longer breaks increase the likelihood that a break indicator that is not perfectly coincident with the break date will appear sig-
nificant, and we can expect the retention to be equal to the nominal significance level outside a $t = T_1 \pm L$ interval.

As before we consider the special case of volcanic functions and also provide results from step shifts for comparison. Figure 5 shows the analytical as well as simulated non-centrality and power around a true break date at $t = 26$ of length $L = 3$ for $\alpha = 0.05$. The Monte Carlo simulations match the theoretical powers and non-centralities closely.

For no break, the analytical power is uniform and equal to the nominal significance level. When there is a break outside of the interval $T_1 \pm L$ the expected retention of the break indicator equals the nominal significance level. For a step shift of a forced length, given (24), the non-centrality decreases linearly as the numerator falls by $1/L$ per shifted period relative to the correct break date. For longer breaks this implies that the power around the true break date is close to uniform. In the case of volcanic functions, due to the particular functional form, the power and retention probability drop more rapidly and peak clearly around the true break date. The special case presented here only considers the properties of a single time-mis-specified indicator of a fixed length in the model. However, model selection in the indicator saturation approach alleviates many of these concerns in practice. When selecting from a full set of break functions (see section 2.1.2) it is less likely that a break function at $T_1 \pm K$ appears significant because the correct $T_1$ indicator is included in the same model, a mis-timed indicator in a fully saturated model would likely appear significant only if a chance draw of the error offsets the shift.

2.2 Properties of indicator functions under the null of no breaks

Under the null hypothesis when there are no breaks in the DGP, there are two primary concerns regarding the inclusion of a full set of break functions in the statistical model. First, when including a full set of break functions, break indicators may be retained spuriously, and second, there may be concerns about the effect on the distributions of coefficients on variables that are known to be relevant – in other words, does saturating a model with irrelevant variables affects relevant ones?

First, we consider the spurious retention of break indicators. Under the null of no breaks, $\lambda = 0$, the DGP from (8) is given by:

$$y = \epsilon$$  \hspace{1cm} (25)

Based on the above results, when using a split-half approach with a full set of break indicators, the expectation of the estimated coefficients in the first half is given by:

$$E[\hat{\gamma}_{(1)}] = E \left[ (D_1' D_1)^{-1} D_1' \epsilon \right] = 0$$  \hspace{1cm} (26)

The same result generalizes to the union of retained indicators $D_U$. Thus, the $t$-statistics of the included break functions will be centred around zero in expectation when there is no break. Using the selection rule that retains the break function $d_j$ if $|t_{d_j}| > c_\alpha$, then $\alpha T/2$ indicators will be retained on average in each half. Combining the retained
**Figure 5:** Power and retention frequency around the break date where the timing of the break functions is imposed without selection: Simulated data with and without shifts (top), associated non-centrality and simulated t-statistics (middle), analytical and simulated power (bottom) around break $\lambda = -10$ at $T = 26$ of length $L = 3$ and interval $T \pm K$ for $\alpha = 0.05$. Left shows no break, middle a step-break, and right panel a volcanic function break. Analytical non-centralities and power are shown in yellow, simulated t-statistics and retention are shown in blue. Dashed lines mark the break occurrence. Outside of an interval $T = 26 \pm L$ the retention probability and analytical power are equal to the nominal significance level $= \alpha = 0.05$. 
indicators in the final set, $\alpha T$ indicators are retained in expectation. The proportion of spurious indicators can thus be controlled through the nominal significance level of selection. The properties under the null are confirmed below using Monte-Carlo simulations.

Table 3 and Figure 6 report the simulation results when there are no breaks in the DGP but a full set of break functions (of the form of volcanic functions) is included. When using a split-half approach with a 1-cut variable selection decision based on the absolute $t$-statistic, the proportion of irrelevant retained indicators is close to the nominal significance level. In practice, when using a multi-path, multi-split procedure (here implemented through Autometrics) the gauge is close to the nominal significance level for low levels of $\alpha$. A conservative approach (low $\alpha \leq 1\%$) is recommended in practice. When compared to results in Castle, Doornik, Hendry, & Pretis 2015, there is little notable difference between different specifications of break functions, consistent with the analytical properties of irrelevant indicators.

<table>
<thead>
<tr>
<th>Significance Level</th>
<th>Split-Half One-Cut</th>
<th>Multi-Path Search</th>
</tr>
</thead>
<tbody>
<tr>
<td>$\alpha = 0.05$</td>
<td>0.056</td>
<td>0.054</td>
</tr>
<tr>
<td>$\alpha = 0.01$</td>
<td>0.013</td>
<td>0.012</td>
</tr>
<tr>
<td>$\alpha = 0.005$</td>
<td>0.007</td>
<td>0.007</td>
</tr>
<tr>
<td>$\alpha = 0.0025$</td>
<td>0.004</td>
<td>0.004</td>
</tr>
<tr>
<td>$\alpha = 0.001$</td>
<td>0.002</td>
<td>0.002</td>
</tr>
</tbody>
</table>

We now assess the second consideration, which is the effect of including a full set of break indicators when theory variables $X$ are included in the model but are not selected over ('forced'). These could include contemporaneous covariates or autoregressive dynamic variables. For the specific case when the elements of $D$ are specified to be impulse indicators, Johansen & Nielsen (2009) derive the asymptotic distribution of $\beta$ in the full split-half approach in stationary and unit-root non-stationary regressions using the equivalence of IIS and one step Huber-skip M estimators. For an iterated procedure (e.g. resembling the multi-block approach in Autometrics) the distributional results under the null for IIS are derived in Johansen & Nielsen (2013). For the general form of designed indicator functions, we follow theory for the sub-steps of split-half estimation where $N \ll T$ in each step, and appeal to simulation results for the overall algorithm. Consider a simple DGP:

$$y = X\beta + \epsilon$$  \hspace{1cm} (27)

where $\epsilon \sim \text{iid}(0,\sigma^2_\epsilon)$ and the elements of $X$ (dynamic or static) are assumed to be

\footnote{Results of high gauge for high significance levels (e.g. $\alpha \geq 0.05$) are consistent with previous results found by Bergamelli & Urga (2013) for step-functions. Once a large number of spurious breaks is retained, it becomes more likely to keep additional spurious breaks. The results for the gauge in Table 3 are consistent with the distributional theory for the gauge in Johansen & Nielsen 2016.}
Figure 6: Simulation results under the null of no break. Left: Proportion of irrelevant retained break functions (gauge) using split-half and multi-path selection for varying $\alpha$ when there is no break. Middle and Right: Simulated distributions and densities of coefficient $\hat{\beta}$ (true $\beta = 0.5$) on forced parameter $x_t$: with (yellow) – and without (blue) – a full set of break functions.

relevant and not selected over. The model relying on the split-half approach saturated with the first half of the break functions is then:

$$y = X\beta + D_1\gamma(1) + \nu$$  \hspace{1cm} (28)

where the true $\gamma(1) = 0$. Following Hendry & Johansen (2015), given that there is no break in the DGP, the inclusion of a full set of irrelevant additional variables $D_1$ need not affect the distribution of the included relevant parameters $\beta$. Orthogonalizing $X$ and $D_1$ by regressing each column of $D_1$ on $X$ yields the estimator $\hat{\beta}^*$ with asymptotic distribution:\n
$$\sqrt{T} \left( \tilde{\beta}^* - \beta \right) \overset{D}{\to} N \left[ \begin{pmatrix} 0 \\ 0 \end{pmatrix}, \sigma^2 \begin{pmatrix} \Sigma_{XX}^{-1} & 0 \\ 0 & \Sigma_{D_1D_1|X}^{-1} \end{pmatrix} \right]$$  \hspace{1cm} (29)

The distribution of the parameters $\hat{\beta}^*$ on the correct variables $X$ is unaffected by the inclusion of the orthogonalized break indicators $D_1$ when there is no break. The equivalent result holds when the second half of break indicators $D_2$ is included and the resulting union of retained indicators from $D_1$ and $D_2$ given that $N < T$. Orthogonalization relative to shifts, however, is not necessary for estimation in practice. Figure 6 shows the simulated distribution of $\hat{\beta}$ for a single $x_t$ when a full set of break functions is included and selected at $\alpha = 0.005$ (yellow) and when break functions are not

\textsuperscript{13}Where $T^{-1}X'X \overset{P}{\to} \Sigma_{XX}$ for stochastic $X$, and $D_1$ is scaled such that either the break length scales with the sample size, or alternatively the break magnitude increases such that $T^{-1}D_1D_1 \rightarrow \Sigma_{D_1}$ is constant, and for stochastic $X$ it holds that $T^{-1}(\tilde{u}'\tilde{u}) \overset{P}{\to} \Sigma_{D_1D_1|X}$ where $\tilde{u} = D_1 - X\hat{\Gamma}$ from orthogonalization regressions. See supplementary material 5.6 for proof based on Hendry & Johansen (2015).
included (blue). The distribution of $\hat{\beta}$ is unaffected by the saturation of a full set of break functions. In practice the main risk is the spurious retention of break indicators, but this can be controlled through a conservative selection mechanism (low $\alpha$).

3 Empirical illustration: Detecting volcanic eruptions in simulated surface air temperature data

Large volcanic eruptions that inject significant amounts of sulphate aerosols into the stratosphere cause short-lived (multi-year) radiative imbalances that induce surface cooling. Over the course of the last several millennia there have been numerous eruptions that have had impacts on global mean temperatures. Identifying their climatic fingerprint is an important scientific endeavour that relies critically on the robust characterization of the timing and magnitude of past volcanism. An accurate understanding of the impact of past eruptions can lead to more accurate estimates of the effect of stratospheric aerosols – to guide policy from geo-engineering to pollution controls. Records of climatically relevant events primarily rely on sulphur deposits in ice cores (see e.g., Gao et al. 2008, and Crowley & Unterman 2012). However, there remains uncertainty in the precise timing, magnitude, and climatic impact of past volcanic activity (Brohan et al. 2012, Mann et al. 2012, Anchukaitis et al. 2012, G. Schmidt et al. 2011, Baillie & McAneney 2015). Statistical methods such as the break detection methodology presented herein can therefore augment previous volcanic reconstruction estimates by providing additional characterizations of the timing and magnitude of temperature responses to volcanic eruptions when coupled with large-scale proxy estimates of past temperature variability e.g., from tree-rings. As a synthetic evaluation of the performance of the break-indicator saturation method, we search for volcanic eruptions in surface air temperature output from model simulations. While there is some disagreement on the timing, magnitude, and climatic impact of real eruptions over the past several millennia, the present simulation is forced with deterministic (known, imposed) eruptions. It therefore can function as a useful tool for assessing the detection efficacy of the proposed statistical methodology in real-world scenarios when the timing and exact DGP of volcanic eruptions are uncertain.

For our empirical illustration we use the Northern Hemisphere (NH) mean surface air temperature from the historical simulation of the National Center for Atmospheric Research (NCAR) Community Climate System Model 4 (CCSM4) and the Last Millennium (LM) simulation (Landrum et al. 2013). These simulations were made available as part of the Coupled and Paleoclimate Model Intercomparison Projects Phases 5 and 3 (CMIP5/PMIP3), respectively (Taylor et al. 2012). Collectively, the two simulations span the period 850-2005 C.E. To imitate potential proxy reconstructions (e.g., tree-ring based), temperatures for extratropical land areas ($30^\circ$–$90^\circ$ N) were extracted from the model and only summer months (June-August) were used to build annual averages. This time period is expected to show the strongest cooling in response to an eruption (e.g., Zanchettin et al. 2013 argue for a winter-warming effect) and is associated with the seasonal sampling window of many proxies such
as dendroclimatic records. Temperatures are reported as anomalies relative to the 1850-1999 mean. The model is forced with the volcanic reconstruction by Gao et al. (2008) that reports volcanic activity as stratospheric sulphate loadings in teragrams (Tg). While the model is forced with multiple radiative forcing conditions (e.g., solar irradiance, greenhouse gases, volcanoes, land cover changes, and anthropogenic aerosol changes), for the present experiments we treat these as unknown and work with the univariate NH mean temperature series, although multivariate models with more forcing variables could improve the detection algorithm. For a real-world scenario, however, estimates of climate-forcing and -sensitivity are uncertain (IPCC 2013) and may prove to be of limited use in explaining non-volcanic temperature variation in proxy reconstructions, particularly in the presence of changes in measurement (see e.g. Pretis & Hendry 2013).

3.1 Simulation design

We design a break function to capture the temperature response to a large-scale volcanic eruption using a simple zero-dimensional energy balance model (EBM) that equates incoming to outgoing energy derived from simple physics-based models of climate (see e.g., section 1 in Rypdal 2012, section 1 in Schwartz 2012, or Pretis 2015a for linking system energy balance models to econometric system models):

\[
C \frac{dT'}{dt} = F - \theta T'
\]  

(30)

where \(\theta\) is the climate feedback, \(C\) is the heat capacity, \(T'\) the temperature deviation from steady state (similar to the measured temperature anomaly as a departure from a long-term average) and \(F\) denotes radiative forcing (the variable that in our system describes the volcanic shock). The feedback response time of the model is given by \(\tau = \frac{C}{\theta}\). Assuming a volcanic forcing effect of an impulse injection of stratospheric aerosols of \(F\) decaying exponentially at rate \(-1/\gamma\) yields the following functional form of a volcanic function for the associated temperature response:

\[
T'_t = d_t = \begin{cases} 
1 & t \leq L \\
0 & t > L \\
\frac{1}{C} e^{-\frac{t}{\tau}} F \left( \frac{\theta}{C} - \frac{1}{\gamma} \right)^{-1} \left[ e^\left(\frac{\theta}{\gamma} - \frac{1}{\gamma} \right) - 1 \right] & t < T_1, t > L 
\end{cases}
\]  

(31)

On visual inspection (see Figure 7) these calibrations closely match the average-model

---

\(^{14}\text{See supplementary material 5.7 for a derivation. For break detection, the function is normalized to sum to 1 over } L.\)
response based on a superposed epoch analysis of all large-scale volcanic eruptions in the climate model (Mass & Portman 1989). The average model response in temperature is a drop by approximately 1–1.5 degrees C, followed by a smooth reversion to the previous mean over a 3-4 year period. While Gao et al. (2008) estimate the retention time for sulphate aerosols to be 2-3 years, a climatic perturbation of 4 years is in line with findings by Landrum et al. (2013). It is important to emphasize that the in-sample response to a volcanic eruption is not used to design the break function – the method is not trained and evaluated on the same set of observations.

In a more theoretical approach, which avoids particular shape parameters, a single peak (impulse) could be followed by auto-regressive reversion to the mean where we search over a full set of impulses and full set of breaking autoregressive coefficients. The DGP for the response variable NH temperature ($T_t$) is:

$$T_t = f(X_t, V_t) + \epsilon_t$$  \hspace{1cm} (32)

To simulate sampling uncertainty of a proxy-based reconstruction we generate 100 replications of the outcome by adding $\epsilon_t \sim \text{IN}(0, \sigma^2)\) to the NH mean temperature. The main results here are presented for simulations setting $\sigma = 0.2$ which is half the sample standard deviation of the NH time-series of 0.4: the effect of the magnitude of noise is explored in Figure 9. The function $f(X_t, V_t)$ mapping volcanic, $V_t$, and other
forcing, $X_t$, on to temperature is unknown, and the observed forcing variables $V_t$ and $X_t$ are equally treated as unknown. As a proof of concept we consider two models (intercept-only, and AR(1) with intercept)\textsuperscript{15} to detect eruptions:

\begin{align*}
  y_t &= \mu + \gamma' d_t + v_t \\
  y_t &= \rho y_{t-1} + \mu + \gamma' d_t + v_t
\end{align*}

where $d_t$ is a full set of volcanic break functions (31) to be selected over.\textsuperscript{16} To reduce computational requirements due to the varying simulation setup, the full-sample is split into 10 sub-samples of $T = 115$ observations each.\textsuperscript{17} There is little difference between full-sample and sub-sampling performance aside from computational speed (supplementary material 5.8 provides the results for a full-sample simulation). Selection is conducted at $\alpha = 0.01$ implying an expected gauge of 1\% (approximately 1 break function spuriously retained per subsample). Higher retention of break functions can be an indicator of model misspecification. Simulations are evaluated based on the retention frequency of known individual volcanic events (potency), the average potency over all volcanoes and the proportion of spurious eruptions detected (gauge).

### 3.2 Empirical illustration results

Figure 8 and Tables 4 and 5 show the results of detected volcanic events in 100 replications of the modelled NH mean temperature\textsuperscript{18} using the model (a) volcanic function. The retained volcanic breaks coincide predominantly with the simulated volcanic eruptions. Few spurious volcanoes are detected, and those that are spurious exhibit retention frequencies drastically lower than those of volcanoes used to force the model.

Most large-scale simulated volcanic eruptions are detected consistently: 74\% of all larger (> 20 Tg) NH eruptions are detected on average within an interval of ±1 year (57\% of all global eruptions, many of which appear to have had little impact.

\textsuperscript{15}Unless otherwise stated, results refer to the intercept-only case. The intercept term and the auto-regressive terms are not selected over.

\textsuperscript{16}Given the specification of the volcanic break function and if $\sigma_\epsilon$ was the only noise added to the DGP, then the approximate expected non-centrality for a single unknown break using (17) is $\lambda \left(0.2 \sqrt{3.7}\right)^{-3} \approx 0.4^{-1}\lambda$ where $\lambda$ is the full temperature response following a volcanic eruption. Since the specified volcanic function has an approximate trough of 0.58 $\lambda$, a temperature drop of 1 degree after a volcanic eruption implies that overall $\lambda \approx 1.7$. Thus in absence of additional noise and for a single volcanic break with an immediate temperature response of 1 degree, the expected $t$-statistic is approximately $\approx 4.3$. The analytical probability of detecting this eruption is roughly: $P(|t| > c_\alpha) \approx 0.96$ for $\alpha = 0.01$. Large eruptions should be consistently detected if the break function is correctly specified and if $\sigma_\epsilon$ was the only source of noise.

\textsuperscript{17}The total sample size is $T = 1155$, resulting in 9 sub-samples of $T = 115$ observations and one sub-sample of $T = 120$ observations. Significance levels are scaled accordingly. Using a 3Ghz processor the sub-sample approach requires $\approx 5$ seconds to cover the entire sample for one replication (across 10 subsamples), compared to $\approx 5$ minutes for one replication using a full-sample approach.

\textsuperscript{18}Retained volcanic functions with positive coefficients are dropped since these likely constitute positive outliers. The focus here lies on the detection of volcanic events which have a negative temperature response.
Figure 8: Detected Model Volcanic Eruptions from 850-2005. Detected (red) volcanic eruptions in the model temperature series from 850-2005 using function (a) modelling a single-period drop followed by a reversion to the mean together with an intercept. Bar height indicates detection frequency [0, 100%] across 100 simulations. Stacked sulphur deposition record used to force model temperatures are shown for Northern hemisphere (blue) and global measurements (orange) in Tg. Simulated model mean temperature anomalies used to detect the above volcanic eruptions are shown in grey. Mean NH surface temperature data is taken from the Last Millenium and historical simulation of the NCAR CCSM4 model as part of the CMIP5/PMIP3 data archive.
on NH temperatures). Consistent with the basic analytical results presented in the previous section, the intervals of selection around the correct break dates are small. While increasing the band from 0 to 1 generally yields an increase in potency, outside of ±1 year there is little difference (see Table 4). An uncertainty in break dates of +/- 1 year can be the result of a monthly dated volcanic forcing record coupled with an annually dated temperature record, e.g. a December eruption will mainly affect the following year. The season of sulphure injection – before or after summer – can cause offsets in the timing of the temperature response. Equally there may be regional sampling biases based on the construction of the NH mean surface air temperature.

Augmenting the designed break functions (a) by an autoregressive model results in nearly similar potency and gauge relative to the baseline model using just a constant (see Table 4 and Figure 9).

The retention frequency of volcanic functions increases with the magnitude of sulphate emissions of the volcanic eruption (Figure 9). While the overall potency for all volcanoes in the Northern Hemisphere within a 1-year interval is 33%, this increases to 74% when larger volcanic eruptions over 20Tg are considered. Given that potency covers all of the volcanic forcing, much of which is small in magnitude, the result is unsurprising. In particular, the lower potency for small eruptions is not driven by an inconsistency in selection of the same volcano over multiple experiments, but rather in the variation in temperature response between volcanoes. Eruptions in 1641 (Parker) and 1600 (Huaynaputina) are detected 100% of the time while the eruption of 1783 (Laki) is not detected in any of the outcomes. In contrast to most of the other volcanoes, Laki is a high-latitude volcano. Because the CCSM4 model uses spatially resolved sulphate estimates, this eruption only affects the northernmost areas and causes only a minor hemispheric cooling of -0.15 degrees, which is much lower in magnitude than that of any of the other major volcanic events (see Figure 7).

Equally, the potency is affected by the chosen standard deviation of the noise process added to the model mean. The main results here are reported for added noise with a standard deviation of half the sample standard deviation. Figure 9 shows the potency for varying levels of noise.

The proportion of spuriously detected volcanoes (gauge) at around 0.02 is close to the nominal significance level (1/T ≈ 0.01). The fact that it is slightly higher is likely due to the misspecification of the model, which is only run on a constant (including an auto-regressive term in the alternate specification) and set of break functions. Any variability in temperature other than volcanic eruptions may be spuriously attributed to the shape of the volcanic functions. This could be controlled by augmenting the model with additional dynamics (e.g. further autoregressive terms, long-term fluctuations through sine-cosine processes) or known forcing series.

Results for volcanic functions (b) are reported in the supplementary material 5.9 (see Tables 9 and 8). Volcanic functions (b) that capture the slower initial decline in temperature yield a slightly higher potency when measured at the precise timing (see Table 9 and Figure 9).

---

19 There is considerable uncertainty on the impact of the Laki eruption, for example A. Schmidt et al. (2012) find the observed NH peak temperature response to Laki to be around -1 degree, suggesting that the last-millennium simulation used here may not reflect the entire impact of the eruption. Notably, the eruption’s noxious fumes at the time were discussed in White’s (1789) treatment of phenology.
Figure 9: Left: detection potency of NH eruptions for given minimum sulphate emissions and timing for functions (a), (a) + AR(1), and (b) at the precise timing $T$ (dashed) and in the interval of $T \pm 1$ (solid). Right: Detection for varying levels of noise added in the simulation for function (a) for all eruptions (dashed) and large eruptions over 20Tg (solid).

Figure 9. Potency for $t = T_i$ for all $i$ NH volcanoes using (b) is 0.32 vs 0.17 for (a) (0.23 vs 0.11 for Global). This result stems from the single drop in function (a) often being most significant in the second period after an eruption if the cooling lasts for two periods. Once we consider the interval of $T_i \pm 1$ years or volcanoes of larger scale the results are nearly identical for functions (a) and (b). Differentiation between one or two-period cooling following an eruption, and thereby further improvements in detection, could be implemented by searching over functions of type (a) and (b) simultaneously controlling the gauge appropriately.

In summary, large-scale volcanic eruptions can consistently be detected within a $\pm 1$ year interval. Even though the model is likely mis-specified when using only a constant, few spurious volcanic eruptions are retained. The signal to noise ratio remains, however, crucial in detection. When the method is applied to real world proxy reconstructions where lower temperature spikes and higher noise levels can be expected, a well specified baseline model for the temperature process will be required against which volcanic events can be detected to ensure a high power of detection.

3.2.1 Forecasting through breaks

While breaks (such as volcanic eruptions) are by their nature stochastic, using a deterministic approach through a full set of break functions allows us to account for the underlying breaks and model the responses deterministically. This can improve forecasts during breaks if the break function is well specified. Once the break is observed (in this case a volcanic eruption), a forecasting model can be augmented with a break indicator where the magnitude is determined through estimation in the first break period. This indicator then acts as a continuous intercept correction, thereby improving the forecast performance during the break. To illustrate this concept, Figure 10 shows a 1-step forecast for NH model mean temperatures following the simulated
Figure 10: 1-Step forecasts through volcanic eruptions using break indicators. Left: Forecast performance across different methods: model mean temperature during the simulated 1641 eruption (yellow), 1-step forecasts from 1641 onwards are shown for using an AR(1) model with volcanic indicator (red), an AR(1) model without a volcanic indicator (blue) and a robust AR(1) forecast (green) (Clements & Hendry 1999). Models are estimated from 1605 until 1641. Right: 1-step forecast root-mean-squared-error over all NH model volcanic eruptions (> 20Tg) for an AR(1) model with volcanic indicator (red), without (blue), and robust AR(1) forecast (green). Using volcanic indicators, on average, improves the forecast performance during the break period. However, when no break occurs (little to no temperature response), using a break indicator can result in higher RMSE as seen for example for the 1783 model Laki eruption.

1641 eruption, together with the root-mean-squared (RMSE) forecast errors for all NH (> 20Tg) model eruptions based on volcanic function (a). Using volcanic indicators to forecast through the breaks yields on average a lower forecast RMSE (RMSE = 0.51) when compared to a simple AR(1) model (RMSE = 0.71) or even a robust forecasting device (RMSE = 0.66) (Clements & Hendry 1999). Crucially this depends on the correct specification of the break function – for volcanic eruptions further improvements could be achieved by switching to volcanic function (b) if the initial cooling lasts for two periods. Detection of breaks based on theory-informed break functions can therefore act as a robust forecasting device through a continuous intercept correction from climate to economic time series.

4 Conclusions

Saturating a regression model with a full set of designed break functions, and removing all but significant ones through a general-to-specific algorithm yields unbiased estimates of the break magnitude and time. By initializing the model with a full set of break functions many of the shortcomings associated with a forward selection or

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20 The robust forecasting device is based on first differences using the forecasting model for $T + 1 | T$ given by: $y_{T+1 | T} = y_T + \hat{\rho} \Delta y_T$ where $\hat{\rho}$ is estimated using an AR(1) model. No error bars are shown on the green robust forecast in Figure 10 due to the non-standard distribution of the forecast.
specific-to-general approach in break detection can be avoided. Analytical properties and non-centralities can be derived for any deterministic break function and can be extended to breaks in random variables when interacted with the deterministic break specifications. The break detection procedure exhibits desirable properties both in the presence of breaks (stable potency across multiple breaks) and under the null hypothesis of no breaks where the spurious retention of break functions can be controlled through a chosen significance level of selection. The multi-path algorithm (Autometrics) outperforms shrinkage-based estimators, especially when facing multiple breaks. We provide some initial insight into uncertainty on the break date by assessing the retention probability of mis-timed break estimators. Break indicator saturation appears to be effective for detecting large-scale temperature responses to volcanic eruptions. This was shown using surface air temperature output from a combined last millennium and historical climate simulation. Statistically searching over a set of break functions consistently detects large eruptions from the simulated surface air temperatures without prior knowledge of their occurrence. This holds promise for future volcanic detection efforts using real-world proxy reconstructions of temperature variability over the last several millennia. More broadly, break detection using designed functions and indicator saturation provide a framework to analyse the detection of breaks of any designed shape at any point in time, with applications ranging from the detection of previously unknown events (such as shifts in time series due to measurement changes or policy impacts), to acting as a robust forecasting device during breaks – from economic recessions to volcanic eruptions.
Table 4: Potency and Gauge for Volcanic Functions (a)

<table>
<thead>
<tr>
<th>Function (a)</th>
<th>$T$</th>
<th>$t = T \pm 1$</th>
<th>$t = T \pm 2$</th>
<th>$t = T \pm 3$</th>
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Gauge NH 0.02
Gauge Global 0.02

Function (a) + AR(1)

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Gauge NH 0.02
Gauge Global 0.02
Table 5: Potency of detection of Volcanic Eruptions $> 20Tg$ using volcanic functions (a) for intervals $t = T \pm 1, 2, 3$

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References


Elliott, G., & Müller, U. K. (2007). Confidence sets for the date of a single break in


5 Appendix

5.1 Power for a known break

The DGP is:

\[ y_t = \mu + \lambda d_t + \epsilon_t \]  \hspace{1cm} (35)

The break shifts \( \mu \) to \( \mu + \lambda d_t \) where \( d_t \) is a break function of length \( L \) beginning at time \( t = T_1 \) where \( T_1 + L \leq T \) such that \( d_t \neq 0 \) for \( T_1 \leq t < T_1 + L \) and 0 otherwise. The correctly specified model is:

\[ y_t = \mu + \gamma d_t + \epsilon_t \]  \hspace{1cm} (36)

Expressions for the estimators \( \hat{\mu}, \hat{\gamma} \) are given by:

\[
\begin{align*}
\left( \hat{\mu} - \mu, \hat{\gamma} - \lambda \right) & = \left( \frac{T}{\sum_{t=T_1}^{T_1+L-1} d_t} \sum_{t=T_1}^{T_1+L-1} d_t^2 \right)^{-1} \left( \frac{\sum_{t=0}^{T} \epsilon_t}{\sum_{t=T_1}^{T_1+L-1} d_t \epsilon_t} \right) \\
& = T^{-1} \left[ \sum_{t=T_1}^{T_1+L-1} d_t^2 - \left( \sum_{t=T_1}^{T_1+L-1} d_t \right)^2 \right] \times \left( \sum_{t=T_1}^{T_1+L-1} d_t \sum_{t=0}^{T} \epsilon_t - \sum_{t=T_1}^{T_1+L-1} d_t \sum_{t=T_1}^{T_1+L-1} d_t \epsilon_t \right)
\end{align*}
\]

\[
\left( \hat{\gamma}_1 \right) = \lambda r + (D'_1 D_1)^{-1} D'_1 \epsilon.
\]

5.2 Proof of equation (10)

Proof that \( \hat{\gamma}(1) = \lambda r + (D'_1 D_1)^{-1} D'_1 \epsilon \).

\[
\hat{\gamma}(1) = \lambda (D'_1 D_1)^{-1} D'_1 d_{T_1} + (D'_1 D_1)^{-1} D'_1 \epsilon
\]

\[
= \lambda (D'_1 D_1)^{-1} (D'_1 D_1) r + (D'_1 D_1)^{-1} D'_1 \epsilon = \lambda r + (D'_1 D_1)^{-1} D'_1 \epsilon
\]

using:

\[
(D'_1 D_1) r = D'_1 \begin{pmatrix} d_1 & \cdots & d_{T_1} & \cdots & d_{T/2} \end{pmatrix} \begin{pmatrix} 1_{T_1} \\ \vdots \\ 0 \end{pmatrix} = D'_1 d_{T_1}
\]

Assuming that the retained indicators of the first half \( D_{1*} \) include the true break indicator, the same result applies to the estimator \( \hat{\gamma}(U) \) over the retained indicators \( D_U = [D_{1*} D_{2*}] \) where \( \hat{\gamma}(U) = \lambda r + (D'_U D_U)^{-1} D'_U \epsilon. \)
5.3 Proof of generalization of step-functions for a known break

Proof that \[ \hat{\mu} - \mu = \bar{\epsilon}_2 \] and \[ \hat{\gamma} - \lambda = \bar{\epsilon}_1 - \bar{\epsilon}_2 \] when \( d_t \) take the form of SIS-type step indicators and a single break from \( t = 0 \) to \( T_1 \) is considered. Note, under these specifications \[ \sum_{t=T_1}^{T_1+L} d_t^2 = \sum_{t=T_1}^{T_1} d_t = T_1. \]

Then
\[
\hat{\mu} - \mu = \frac{T_1 \left( \sum_{t=1}^{T} \epsilon_t - \sum_{t=1}^{T_1} \epsilon_t \right)}{T (T_1 - T_1^2 / T)} = \frac{\sum_{t=T_1+1}^{T} \epsilon_t}{T - T_1} = \bar{\epsilon}_2
\]

and
\[
\hat{\gamma} - \lambda = \frac{T \sum_{t=1}^{T_1} \epsilon_t - T_1 \sum_{t=1}^{T} \epsilon_t}{T_1 (T - T_1)} = \frac{T_1 (T - T_1) \bar{\epsilon}_1 - T_1 (T - T_1) \bar{\epsilon}_2}{T_1 (T - T_1)} = \bar{\epsilon}_1 - \bar{\epsilon}_2
\]

using \( \bar{\epsilon}_1 = 1/T_1 \sum_{t=1}^{T_1} \epsilon_t \) and \( \bar{\epsilon}_2 = 1/(T - T_1) \sum_{t=T_1+1}^{T} \epsilon_t \).

5.4 Algorithm specifications

Here we outline the split-half algorithm (Algorithm 1, see Figure 1) used for theory-derivations and simulations, and sketch the multi-path algorithm (Algorithm 2) used in simulations and in practice:

**Algorithm 1: Split-Half Indicator Saturation for Designed Break Functions.** Choose a target level of significance \( \alpha \).

1. Design a break function and construct the \((T \times T)\) break matrix \( D \) using equation (4).
2. Split the break matrix \( D \) into two sets: \( D_1 = (d_1, \ldots, d_{T/2}) \) and \( D_2 = (d_{T/2+1}, \ldots, d_T) \).
3. Compute the least squares estimators \( \hat{\gamma}_{(1)} \) and \( \hat{\gamma}_{(2)} \) using (10) and (12) fixing an intercept (and additional covariates if required) in each regression.
4. Construct the union set of \( k \) retained break functions as \( D_U = (d_j, \ldots, d_{j+k}) \) where \( d_j \) is included if \( |\hat{t}_{\gamma_{i,j}}| \geq c_\alpha \) in step 3, where \( i = 1, 2 \) for each set.
5. Compute the least squares estimator \( \hat{\gamma}_{(U)} \) using (14), fixing an intercept (and additional covariates if required) in each regression. Retain final break functions \( j \) if \( |\hat{t}_{\gamma_{U,j}}| \geq c_\alpha \).
Algorithm 2: Multi-Path Indicator Saturation for Designed Break Functions.

1. Design a break function and construct the \((T \times T)\) break matrix \(D\) using equation (4).

2. Use a multi-path and multi-block algorithm: for the present simulations and application we rely on the parallel step-wise backwards tree-search algorithm Auto-metrics (Doornik 2009a) fixing an intercept (and additional covariates if required) in the general unrestricted model (2).

3. Retain the break functions from the terminal model chosen by the specified tiebreaker criterion.

5.5 Simulation results for a simple dynamic autoregressive DGP

Here we present the results on power and false-positive rate when using a multi-path search and the DGP includes an autoregressive term of order one. The DGP is given by:

\[
y_t = \rho y_{t-1} + \lambda d_t + \epsilon_t
\]

where \(\epsilon_t \sim \text{IN}(0, \sigma^2_\epsilon)\), \(\sigma^2_\epsilon = 1\) and \(\rho = 0.3\). The model includes an AR(1) term and a full set of break functions. Figure 6 and Table 6 provide the results. The null-retention (gauge) remains well calibrated to the nominal size, the power is slightly lower compared to the static case, likely due to the shape of the break mimicking an auto-regressive process and thereby being detected less frequently.

Table 6: Power of detecting an unknown break when multi-path searches in an auto-regressive DGP and model. Statistics were generated from 1000 simulations and detection significance was set to \(\alpha = 0.01\), with a length of \(L = 3\). Break magnitude \(\lambda\) corresponds to the full response over the entire break, the peak is \(0.58\lambda\), T=100.

<table>
<thead>
<tr>
<th>Multi-Path</th>
<th>Potency</th>
<th>Gauge</th>
</tr>
</thead>
<tbody>
<tr>
<td>(\lambda = 6), peak=3.48</td>
<td>0.71</td>
<td>0.018</td>
</tr>
<tr>
<td>(\lambda = 4), peak=2.23</td>
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<td>0.015</td>
</tr>
<tr>
<td>(\lambda = 2), peak=1.16</td>
<td>0.09</td>
<td>0.016</td>
</tr>
</tbody>
</table>

5.6 Proof of null results when variables not-selected-over are included

We investigate the approximate distributional consequences of including a full set of irrelevant break variables into a model with assumed \(K\) relevant variables \(X\) with associated coefficients \(\beta\). This is based on the results of Hendry & Johansen (2015). The DGP is:

\[
y = X\beta + \epsilon
\]
Figure 11: Left: Potency of detecting a volcanic break of magnitude $\lambda$ for level of significance $\alpha$ using multi-path selection for an autoregressive DGP and model. Right: Proportion of spuriously retained break indicators (gauge). Break magnitude $\lambda$ corresponds to the full response over the entire break, the peak is $0.58\lambda$.

where $\epsilon \sim \text{iid}(0, \sigma^2\epsilon I)$. The first-half model is given as:

$$y = X\beta + D_1\gamma^{(1)} + v$$  \hspace{1cm} (46)

where the true $\gamma^{(1)} = 0$. Consider first orthogonalizing $X$ and $D_1$ by regressing each column of $D_1$ on $X$, this yields the $(K \times T/2)$ matrix of coefficients $\hat{\Gamma}$:

$$\hat{\Gamma} = (X'X)^{-1}X'D_1$$  \hspace{1cm} (47)

The $(T \times T/2)$ residuals $\hat{u}$ are defined through:

$$D_1 = X\hat{\Gamma} + \hat{u}$$  \hspace{1cm} (48)

where $\hat{u}'X = 0$. Substituting for $D_1$ in the model equation yields:

$$y = X\beta + (X\hat{\Gamma} + \hat{u})\gamma^{(1)} + v$$  \hspace{1cm} (49)

$$= X\left(\beta + \hat{\Gamma}\gamma^{(1)}\right) + \hat{u}\gamma^{(1)} + v$$  \hspace{1cm} (50)

$$= X\beta^* + \hat{u}\gamma^{(1)} + v$$  \hspace{1cm} (51)
where $\beta^* = (\beta + \hat{\Gamma}(1)) = \beta$ as $\gamma(1) = 0$. Then:

\[
\begin{pmatrix}
\hat{\beta} - \beta \\
\hat{\gamma}(1) - 0
\end{pmatrix} = \begin{pmatrix}
\hat{\Gamma} \\
\hat{u}X
\end{pmatrix}^{-1} \begin{pmatrix}
\epsilon' \\
\hat{\epsilon'}
\end{pmatrix}
\]

(52)

\[
\begin{pmatrix}
\hat{\Gamma} \\
\hat{u}X
\end{pmatrix}^{-1} \begin{pmatrix}
\hat{\epsilon}' \\
\hat{u}'
\end{pmatrix}
\]

(53)

since $\hat{u}'X = 0$. Under the assumption that $T^{-1}X'X \xrightarrow{P} \Sigma_{XX}$, the asymptotic distribution is then:

\[
\sqrt{T} \begin{pmatrix}
\hat{\beta} - \beta \\
\hat{\gamma}(1) - 0
\end{pmatrix} \xrightarrow{D} N \left( \begin{pmatrix} 0 \\ 0 \end{pmatrix}, \sigma^2 \begin{pmatrix}
\Sigma_{XX}^{-1} & 0 \\
0 & \Sigma_{D_1D_1|X}
\end{pmatrix} \right)
\]

(54)

The distribution of the parameters $\beta$ on the correct variables $X$ is unaffected by the inclusion of the break indicators $D_1$ when there is no break. The equivalent result holds when the second half of break indicators $D_2$ is included. In practice the main risk is the spurious retention of break indicators, but this can be controlled through a conservative selection rule.

### 5.7 Energy balance model response to a large volcanic eruption

The basic energy balance model in terms of deviations from the steady state is given by:

\[
C\frac{dT'}{dt} = F(t) - \theta T'
\]

(55)

The differential equation can be solved using an integrating factor. Re-arranging the above model yields:

\[
\frac{dT'}{dt} + \frac{\theta}{C} T' = \frac{1}{C} F(t)
\]

(56)

\[
\frac{dT'}{dt} + \frac{1}{\tau} T' = \frac{1}{C} F(t)
\]

(57)

Using the integrating factor $I(t) = e^{\frac{t}{\tau}}$ yields

\[
\frac{d}{dt} T' e^{\frac{t}{\tau}} = \frac{F(t)}{C} e^{\frac{t}{\tau}}
\]

(58)

\[
T' e^{\frac{t}{\tau}} = \int_0^t \frac{F(u)}{C} e^{\frac{u}{\tau}} du
\]

(59)

\[
T' = \frac{1}{C} e^{\frac{-t}{\tau}} \int_0^t F(u) e^{\frac{u}{\tau}} du
\]

(60)

This expression can be used to solve for the temperature response following a large volcanic eruption under the assumption that the volcanic eruption is approximated by a pulse forcing followed by exponential decay at rate $-1/\gamma$. Let $F(t)$ denote the volcanic forcing:

\[
F(t) = 0 \text{ for } t \leq 0 \text{ and } F(t) = F e^{\frac{-t}{\tau}} \text{ for } t > 0
\]

(61)
Using this expression yields:

\[
T' = \frac{1}{C} e^{t_0^t} \int_0^t F(u) e^{\frac{u}{\tau}} du \\
T' = \frac{1}{C} e^{t_0^t} F \left( \frac{\theta}{C} - \frac{1}{\gamma} \right)^{-1} \left[ e^{t_0^t \tau} - 1 \right]
\] (62) (63)

5.8 Simulation results using a full sample search

Table 7 provides simulation results when a full-sample rather than a sub-sample approach is used for function (a) with an intercept-only model. There is little difference relative to the sub-sampling method at the cost of increased computational time. Using a 3Ghz processor the sub-sample approach requires \( \approx 5 \) seconds to cover the entire sample for one replication (across 10 subsamples), compared to \( \approx 5 \) minutes for one replication using a full-sample approach, similar to the analysis in Doornik & Hendry (2015).

Table 7: Potency and Gauge for Volcanic Functions (a) using a full sample search

<table>
<thead>
<tr>
<th></th>
<th>( T )</th>
<th>( t = T \pm 1 )</th>
<th>( t = T \pm 2 )</th>
<th>( t = T \pm 3 )</th>
</tr>
</thead>
<tbody>
<tr>
<td>Potency NH Tg&gt; 20</td>
<td>0.32</td>
<td>0.67</td>
<td>0.67</td>
<td>0.67</td>
</tr>
<tr>
<td>Potency NH Tg&gt; 0</td>
<td>0.22</td>
<td>0.53</td>
<td>0.57</td>
<td>0.58</td>
</tr>
<tr>
<td>Potency Global Tg&gt;20</td>
<td>0.23</td>
<td>0.52</td>
<td>0.57</td>
<td>0.58</td>
</tr>
<tr>
<td>Potency Global Tg&gt; 0</td>
<td>0.22</td>
<td>0.53</td>
<td>0.57</td>
<td>0.58</td>
</tr>
<tr>
<td>Gauge NH</td>
<td>0.02</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Gauge Global</td>
<td>0.02</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
### 5.9 Results using volcanic function (b)

**Table 8: Potency and Gauge for Volcanic Functions (b)**

<table>
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<tr>
<th></th>
<th>$T$</th>
<th>$t = T \pm 1$</th>
<th>$t = T \pm 2$</th>
<th>$t = T \pm 3$</th>
</tr>
</thead>
<tbody>
<tr>
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<td>0.60</td>
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</tr>
<tr>
<td>Potency NH Tg &gt; 0</td>
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<td>0.32</td>
<td>0.33</td>
<td>0.34</td>
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<td>0.57</td>
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<td>Gauge Global</td>
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Table 9: Detection of Volcanic Eruptions > 20Tg using volcanic functions (b)

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<tr>
<th>NH Volcano</th>
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<th>Potency</th>
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<table>
<thead>
<tr>
<th>Global Volcano</th>
<th>Tg</th>
<th>Potency</th>
<th>t = T</th>
<th>t = T ±1</th>
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