Approximate Cross-Validation for Structured Models

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Abstract

Many modern data analyses benefit from explicitly modeling dependence structure in data – such as measurements across time or space, ordered words in a sentence, or genes in a genome. Cross-validation is the gold standard to evaluate these analyses but can be prohibitively slow due to the need to re-run already-expensive learning algorithms many times. Previous work has shown approximate cross-validation (ACV) methods provide a fast and provably accurate alternative in the setting of empirical risk minimization. But this existing ACV work is restricted to simpler models by the assumptions that (i) data are independent and (ii) an exact initial model fit is available. In structured data analyses, (i) is always untrue, and (ii) is often untrue. In the present work, we address (i) by extending ACV to models with dependence structure. To address (ii), we verify – both theoretically and empirically – that ACV quality deteriorates smoothly with noise in the initial fit. We demonstrate the accuracy and computational benefits of our proposed methods on a diverse set of real-world applications.

1 Introduction

Models with complex dependency structures have become standard machine learning tools in analyses of data from science, social science, and engineering fields. These models are used to characterize disease progression [Sukkar et al., 2012, Wang et al., 2014, Sun et al., 2019], to track crime in a city [Balocchi and Jensen, 2019, Balocchi et al., 2019], and to monitor and potentially manage traffic flow [Ihler et al., 2006, Zheng and Liu, 2017] among many other applications. The potential societal impact of these methods necessitates that they be used and evaluated with care. Indeed, recent work [Musgrave et al., 2020] has emphasized that hyperparameter tuning and assessment with cross-validation (CV) [Stone, 1974, Geisser, 1975] is crucial to trustworthy and meaningful analysis of modern, complex machine learning methods.

While CV offers a conceptually simple and widely used tool for evaluation, it is computationally demanding in complex models; these models often already face severe computational demands to fit just once, and CV requires multiple re-fits. To address these prohibitive computational costs, recent authors [Beirami et al., 2017, Rad and Maleki, 2020, Giordano et al., 2019] have proposed to instead use an approximation to CV (ACV); their work demonstrates that ACV methods perform well in both theory and practice for a collection of practical models. These methods take two principal forms: one

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approximation based on a Newton step (NS) \cite{Beirami2017, Rad2020} and one based on the classical infinitesimal jackknife (IJ) from statistics \cite{Koh2017, Beirami2017, Giordano2019}. Though both ACV forms show promise, there remain major roadblocks to applying either NS or IJ to models with dependency structure. First, all existing ACV theory and algorithms assume that data are independent – an assumption necessarily violated by models that explicitly capture dependency in time, space, or other modalities. E.g. to evaluate time series models, we often need to be able to drop out data points within the time series itself – and similarly for spatiotemporal or other dependency structures. Second, NS methods require recomputation and inversion of a model’s Hessian matrix at each CV fold. In the complex models we consider here, this cost can itself be prohibitive. Finally, existing theory for IJ methods requires an exact initial fit of the model – and authors so far have taken great care to obtain such a fit \cite{Giordano2019, Stephenson2020}. But practitioners learning in e.g. large sequences or graphs typically settle for an approximate fit to limit computational cost.

In this paper, we address these concerns and thereby expand the reach of ACV to include more sophisticated models with dependencies among data points and for which exact model fits are infeasible. To avoid the cost of matrix recomputation and inversion across folds, we here focus on the IJ, rather than the NS. In particular, in Section \ref{sec:ij} we develop IJ approximations for dropping out individual nodes in a dependence graph. Our methods allow us e.g. to leave out points within or at the end of a time series – but our methods also apply to more general Markov random fields, without a strict chain structure. In Section \ref{sec:experiments}, we demonstrate that the IJ yields a useful ACV method even without an exact initial model fit. In fact, we show that the quality of the IJ approximation decays with the quality of the initial fit in a smooth and interpretable manner. Finally, we demonstrate our method on a diverse set of real-world applications and models in Section \ref{sec:results} and Appendix \ref{app:results}. These include count data analysis with time-varying Poisson processes, named entity recognition with neural conditional random fields, motion capture analysis with auto-regressive hidden Markov models, and a spatial analysis of crime data with hidden Markov random fields.

2 Structured models and cross-validation

2.1 Structured models

Throughout we consider two types of models: (1) hidden Markov random fields (MRFs) with observations \(x\) and latent variables \(z\) and (2) conditional random fields (CRFs) with inputs (i.e., covariates) \(x\) and labels \(z\), both observed. Our developments for hidden MRFs and CRFs are very similar, but with slight differences. We detail MRFs in the main text; throughout, we will refer the reader to the appendix for the CRF treatment. We first give an illustrative example of MRFs and then the general formulation; a CRF overview appears in Appendix \ref{app:crf}.

Example: Hidden Markov Models (HMMs) capture sequences of observations such as words in a sentence or longitudinally measured physiological signals. Consider an HMM with \(N\) (independent) sequences, \(T\) time steps, and \(K\) states. We take each observation to have dimension \(R\). So the \(t\)th observed element in the \(n\)th sequence is \(x_{nt} \in \mathbb{R}^R\), and the latent \(z_{nt} \in [K] := \{1, \ldots, K\}\). The model is specified by (1) a distribution on the initial latent \(p(z_{n1}) = \text{Cat}(z_{n1} \mid \pi)\), where \(\text{Cat}\) is the categorical distribution and \(\pi \in \Delta_{K-1}\), the \(K-1\) simplex; (2) a \(K \times K\) transition matrix \(A\) with columns \(A_k \in \Delta_{K-1}\) and \(p(z_{nt} \mid z_{n,t-1}) = \text{Cat}(z_{nt} \mid A_{zt_{t-1}})\); and (3) emission distributions \(F\) with parameters \(\theta_k\) such that \(p(x_{nt} \mid z_{nt}) = F(x_{nt} \mid \theta_{zn_t})\). We collect all parameters of the model in \(\Theta := \{\pi, \{A_k\}_{k=1}^{K}, \{\theta_k\}_{k=1}^{K}\}\). We consider \(\Theta\) as a vector of length \(D\). We may have a prior \(p(\Theta)\).

More generally, we consider (hidden) MRFs with \(N\) structured observations \(x_n\) and latents \(z_n\), independent across \(n \in [N]\). We index single observations of dimension \(R\) (respectively, latents) within the structure by \(t \in [T]\); \(x_{nt} \in \mathbb{R}^R\) (respectively, \(z_{nt}\)). Our experiments will focus on bounded, discrete \(z_{nt}\) (i.e., \(z_{nt} \in [K]\)), but we use more inclusive notation (that might e.g. apply to continuous latents) when possible. We consider models with parameters \(\Theta \in \mathbb{R}^D\) and a single emission factor for each latent.

\[
- \log p(x, z; \Theta) = Z(\Theta) + \sum_{n=1}^{N} \left\{ \sum_{t \in [T]} \psi_t(x_{nt}, z_{nt}; \Theta) + \sum_{c \in F} \phi_c(z_{nc}; \Theta) \right\}, \tag{1}
\]
where $z_{ne} := (z_{nt})_{t \in c}$ for $c \subseteq [T]$; $\psi_\lambda$ is a log factor mapping $(x_{nt}, z_{nt})$ to $\mathbb{R}$; $\phi_\lambda$ is a log factor mapping collections of latents, indexed by $c$, to $\mathbb{R}$; $F$ collects the subsets indexing factors; and $Z(\Theta)$ is a negative log normalizing constant. HMMs, as described above, are a special case; see Appendix [B] for details. For any MRF, we can learn the parameters by marginalizing the latents and maximizing the posterior, or equivalently the joint, in $\Theta$. Maximum likelihood estimation is the special case with formal prior $p(\Theta)$ constant across $\Theta$.

$$\hat{\Theta} := \arg\min_{\Theta} -\log p(x; \Theta) - \log p(\Theta) = \arg\min_{\Theta} \int_{z} -\log p(x; z, \Theta) \, dz - \log p(\Theta). \quad (2)$$

### 2.2 Challenges of cross-validation and approximate cross-validation in structured models

In CV procedures, we iteratively leave out some data in order to diagnose variation in $\hat{\Theta}$ under natural data variability or to estimate the predictive accuracy of our model. We consider two types of CV of interest in structured models; we make these formulations precise later. (1) We say that we consider leave-within-structure-out CV (LWCV) when we remove some data points $x_{nt}$ within a structure and learn on the remaining data points. For instance, we might try to predict crime in certain census tracts based on observations in other tracts. Often in this case $N = 1$ [Celeux and Durand 2008, Hyndman and Athanasopoulos 2018 Chapter 3.4], and we assume LWCV has $N = 1$ for notational simplicity in what follows. (2) We say that we consider leave-structure-out CV (LSCV) when we leave out entire $x_n$, for either a single $n$ or a collection of $n$. For instance, with a state-space model of gene expression, we might predict some individuals’ gene expression profiles given other individuals’ profiles. In this case, $N \gg 1$ [Rangel et al. 2004, DeCaprio et al. 2007]. In either (1) or (2), the goal of CV is to consider multiple folds, or subsets of data, left out to assess variability and improve estimation of held-out error. But every fold incurs the cost of the learning procedure in Eq. (2).

Indeed, practitioners have explicitly noted the high cost of using multiple folds and have resorted to using only a few, large folds [Celeux and Durand 2008], leading to biased or noisy estimates of the out-of-sample variability.

A number of researchers have addressed the prohibitive cost of CV with approximate CV (ACV) procedures for simpler models [Beirami et al. 2017, Rad and Maleki 2020, Giordano et al. 2019]. Existing work focuses on the following learning problem with weights $w \in \mathbb{R}^D$:

$$\hat{\Theta}(w) = \arg\min_{\Theta} \sum_{j \in [J]} w_j f_j(\Theta) + \lambda R(\Theta), \quad (3)$$

where $\forall j \in [J], f_j, R : \mathbb{R}^D \to \mathbb{R}$ and $\lambda \in \mathbb{R}_+$. When the weight vector $w$ equals the all-ones vector $1_J$, we recover an empirical loss minimization problem. By considering all weight vectors with one weight equal to zero, we recover the folds of leave-one-out CV; other forms of CV can be similarly recovered. The notation $\hat{\Theta}(w)$ emphasizes that the learned parameter values depend on the weights.

To see if this framework applies to LWCV or LSCV, we can interpret $f_j$ as a negative log likelihood (up to normalization) for the $j$th data point and $\lambda R$ as a negative log prior. Then the likelihood corresponding to the objective of Eq. (3) factorizes as $p(x \mid \Theta) = \prod_{j \in [J]} p(x_j \mid \Theta) \propto \prod_{j \in [J]} \exp(-f_j(\Theta))$. This factorization amounts to an independence assumption across the $\{x_j\}_{j \in [J]}$. In the case of LWCV, with $N = 1, j$ must serve the role of $t$, and $J = T$. But the $x_t$ are not independent, so we cannot apply existing ACV methods. In the LSCV case, $N \geq 1$, and $j$ in Eq. (5) can be seen as serving the role of $n$, with $J = N$. Since the $x_n$ are independent, Eq. (5) can express LSCV folds.

**Previous ACV work** provides two primary options for the LSCV case. We give a brief review here, but see Appendix [X] for a more detailed review. One option is based on taking a single Newton step on the LSCV objective starting from $\hat{\Theta}(1_T)$ [Beirami et al. 2017, Rad and Maleki 2020]. Except in special cases – such as leave-one-out CV for generalized linear models – this Newton-step approach requires both computing and inverting a new Hessian matrix for each fold, often a prohibitive expense; see Appendix [H] for a discussion. An alternative method [Koh and Liang 2017, Beirami et al. 2017, Giordano et al. 2019] based on the infinitesimal jackknife (IJ) from statistics [Jaekel 1972, Efron 1981] constructs a Taylor expansion of $\hat{\Theta}(w)$ around $w = 1_T$. For any model of the form in Eq. (3), the IJ requires just a single Hessian matrix computation and inversion. Therefore, we focus on the IJ for LSCV and use the IJ for inspiration when developing LWCV below. However, all existing IJ theory and empirics require access to an *exact* minimum for $\hat{\Theta}(1_J)$. Indeed, previous authors [Giordano et al. 2019, Stephenson and Broderick 2020] have taken great care to find an
exact minimum of Eq. (3). Unfortunately, for most complex, structured models with large datasets, finding an exact minimum requires an impractical amount of computation. Others [Bürkner et al., 2020] have developed ACV methods for Bayesian time series models and for Bayesian models without dependence structures [Vehtari et al., 2017]. Our development here focuses on empirical risk minimization and is not restricted to temporal models.

In the following, we extend the reach of ACV beyond LSCV and address the issue of inexact optimization. In Section [1], we adapt the IJ framework to the LWCV problem for structured models. In Section [2], we show theoretically that both our new IJ approximation for LWCV and the existing IJ approximation applied to LSCV are not overly dependent on having an exact optimum. We support both of these results with practical experiments in Section [5].

3 Cross-validation and approximate cross-validation in structured models

We first specify a weighting scheme, analogous to Eq. (3), to describe LWCV in structured models; then we develop ACV for this scheme. Recall that CV in independent models takes various forms such as leave-\( k \)-out and \( k \)-fold CV. Similarly, we consider the possibility of leaving out\( k \) arbitrary sets of data indices \( o \in O \), where each \( o \subseteq [T] \). We have two options for how to leave data out in hidden MRFs; see Appendix [C] for CRFs. (A) For each data index \( t \) left out, we leave out the data point \( x_t \) but we retain the latent \( z_t \). For instance, in a time series, if data is missing in the middle of the series, we still know the time relation between the surrounding points, and would leave in the latent to maintain this relation. (B) For each data index \( t \) left out, we leave out the data point \( x_t \) and the latent \( z_t \). For instance, consider data in the future of a time series or pixels beyond the edge of a picture. We typically would not include the possibility of all possible adjacent latents in a structure, so leaving out \( z_t \) as well is more natural. In either case, analogous to Eq. (3), \( \Theta(w) \) is a function of \( w \) computed by minimizing the negative log joint \(- \log p(x; \Theta, w) - \log p(\Theta)\), now with \( w \) dependence, in \( \Theta \). For case (A), we adapt Eq. (1) (with \( N = 1 \)) and Eq. (2) with a weight \( w_t \) for each \( x_t \) term:

\[
\hat{\Theta}(w) = \arg\min_{\Theta} Z(\Theta, w) + \int_z \left[ \sum_{t \in [T]} w_t \psi_t(x_t, z_t; \Theta) \right] + \left[ \sum_{c \in F} \phi_c(z_c; \Theta) \right] \, dz - \log p(\Theta). \tag{4}
\]

Note that the negative log normalizing constant \( Z(\Theta, w) \) may now depend on \( w \) as well. For case (B), we adapt Eq. (1) and Eq. (2) with a weight \( w_t \) for each term with \( x_t \) or \( z_t \):

\[
\hat{\Theta}(w) = \arg\min_{\Theta} Z(\Theta, w) + \int_z \left[ \sum_{t \in [T]} w_t \psi_t(x_t, z_t; \Theta) \right] + \left[ \sum_{c \in F} \left( \prod_{t \in c} w_t \right) \phi_c(z_c; \Theta) \right] \, dz - \log p(\Theta). \tag{5}
\]

In both cases, the choice \( w = 1_T \) recovers the original learning problem. Likewise, setting \( w = w_o \), where \( w_o \) is a vector of ones with \( w_{t_o} = 0 \) if \( t \in o \), drops out the data points in \( o \) (and latents in case (B)). We show in Appendix [C] that these two schemes are equivalent in the case of chain-structured graphs when \( o = \{T', T' + 1, \ldots, T\} \) but also that they are not equivalent in general. We thus consider both schemes going forward.

The expressions above allow a unifying viewpoint on LWCV but still require re-solving \( \hat{\Theta}(w_o) \) for each new CV fold \( o \). To avoid this expense, we propose to use an IJ approach. In particular, as discussed by Giordano et al., [2019], the intuition of the IJ is to notice that, subject to regularity conditions, a small change in \( w \) induces a small change in \( \hat{\Theta}(w) \). So we propose to approximate \( \hat{\Theta}(w_o) \) with \( \Theta_{ACV}(w_o) \), a first-order Taylor series expansion of \( \hat{\Theta}(w) \) as a function of \( w \) around \( w = 1_T \). We follow Giordano et al., [2019] to derive this expansion in Appendix [1]. We summarize our method and define \( \Theta_{ACV} \), with three arguments, in Algorithm [1]. We define \( \Theta_{ACV}(w_o) := \Theta_{ACV}(\Theta(1_T), x, o) \).

First, note that the proposed procedure applies to either weighting style (A) or (B) above; they each determine a different \( \log p(x; \Theta, w) \) in Algorithm [1]. We provide analogous LSCV algorithms for MRFs and CRFs in Algorithms [2] and [3], Appendices [C] and [D]. Next, we compare the cost of our

\footnote{Note that the weight formulation could be extended to even more general reweightings in the spirit of the bootstrap. Exploring the bootstrap for structured models is outside the scope of the present paper.}
We focus here on a particular use of CV: estimating out-of-sample loss. For simplicity, we discuss the
want
Algorithm 1
\[ \varepsilon \]
(2) The difference in the parameter value, \( \varepsilon \)
(Algorithm 1) and LSCV (Algorithms 2 and 3 in the appendix) assume we have access to the exact
M
(\text{cost } W_2 \text{ and } 3) is in
Proposition 1.
Let \( \varepsilon \)
below, we will bound the error in our approximation:
\[ \left| L_{\text{ACV}}(\Theta_1, x, o) \right| \]
\[ = 1 \]
N
(1)
(2)
\[ \text{input to Algorithm 1; we denote this approximation by} \]
\[ \varepsilon \]
\[ \text{for an extension to CV computed with a generic loss} \]
\[ \ell \]
\[ \text{sequence of parameter values} \]
\[ \Theta \]
By envisioning the IJ as a Taylor series approximation around \( \hat{\Theta}(1_T) \), the approximations for LWCV
Algorithm 1 and LSCV (Algorithms 2 and 3 in the appendix) assume we have access to the exact
optimum \( \hat{\Theta}(1_T) \). In practice, though, especially in complex problems, computational considerations
often require using an inexact optimum. More precisely, any optimization algorithm returns a
\[ \hat{\Theta}(1_T) \]
\[ \text{cost of any one of our ACV algorithms (Algorithms 1, 2 and 3)} \]
is in \( O(MN + (D^2 + |O|) \cdot |O|) \). Exact CV is in \( O(MNS \cdot |O|) \).

\[ \text{Proof.} \] For each of the \( |O| \) folds of CV and each of the \( N \) structures, we compute the marginalization
(cost \( M \)) at each of the \( S \) steps of the optimization procedure. In our ACV algorithms, we compute \( H \)
and \( J \) with automatic differentiation tools [Baydin et al. 2018]. The results of Bartholomew-Biggs et al. [2000]
demonstrate that \( H \) and \( J \) each require the same computation (up to a constant) as WEIGHTEDMARG.
So, across \( N \), we incur cost \( MN \). The remaining cost is from the for loop.

In structured problems, we generally expect \( M \) to be large; see Appendix D for a discussion of the
costs, including in the special case of chain-structured MRFs and CRFs. And for reliable CV, we
want \( |O| \) to be large. So we see that our ACV algorithms reap a savings by, roughly, breaking up the
product of these terms into a sum and avoiding the further \( S \) multiplier.

4 IJ behavior under inexact optimization

By envisioning the IJ as a Taylor series approximation around \( \hat{\Theta}(1_T) \), the approximations for LWCV
(Algorithm 1) and LSCV (Algorithms 2 and 3 in the appendix) assume we have access to the exact
optimum \( \hat{\Theta}(1_T) \). In practice, though, especially in complex problems, computational considerations
often require using an inexact optimum. More precisely, any optimization algorithm returns a
sequence of parameter values \( (\Theta^{(s)})_{s=1}^{S} \). Ideally the values \( \Theta^{(S)} \) will approach the optimum \( \hat{\Theta}(1_T) \)
as \( S \to \infty \). But we often choose \( S \) such that \( \Theta^{(S)} \) is much farther from \( \hat{\Theta}(1_T) \) than machine
precision. In practice, then, we input \( \Theta^{(S)} \) (rather than \( \hat{\Theta}(1_T) \)) to Algorithm 1. We now check that
the error induced by this substitution is acceptably low.

We focus here on a particular use of CV: estimating out-of-sample loss. For simplicity, we discuss the
\[ N = 1 \]
case here; see Appendix D for the \( N \geq 1 \) case. For each fold \( o \in O \), we compute \( \Theta(w_o) \) from the
points kept in and then calculate the loss (in our experiments here, negative log likelihood) on the
left-out points. I.e. the CV estimate of the out-of-sample loss is \( L_{\text{CV}} := \sum_{o \in O} - \log p(x_o | x_{[T]-o}; \Theta(w_o)) \), where \( - \log p \) may come from either weighting scheme (A) or (B). See Appendix D for an extension to CV computed with a generic loss \( \ell \). We approximate \( L_{\text{CV}} \) using some \( \Theta \) as
input to Algorithm 1, we denote this approximation by \( L_{\ell}^{(T)}(\Theta) := \sum_{o \in O} - \log p(x_o | x_{[T]-o}; \Theta_{\text{ACV}}(\Theta, x, o)) \).

Below, we will bound the error in our approximation: \( |L_{\text{CV}} - L_{\ell}^{(T)}(\Theta^{(S)})| \). There are two sources
of error. (1) The difference in loss between exact CV and the exact IJ approximation, \( \varepsilon_{IJ} \) in Eq. (6).
(2) The difference in the parameter value, \( \varepsilon_{\Theta} \) in Eq. (6), which will control the difference between
\[ L_{11}(\hat{\Theta}(1_T)) \text{ and } L_{11}(\Theta^{(S)}). \]

\[ \varepsilon_{11} := |L_{CV} - L_{11}(\hat{\Theta}(1_T))|, \quad \varepsilon_{\Theta} := ||\Theta^{(S)} - \hat{\Theta}(1_T)||_2 \]

(6)

Our bound below will depend on these constants. We observe that empirics, as well as theory based on the Taylor series expansion underlying the IJ, have established that \( \varepsilon_{11} \) is small in various models; we expect the same to hold here. Also, \( \varepsilon_{\Theta} \) should be small for large enough \( S \) according to the guarantees of standard optimization algorithms. We now state some additional regularity assumptions before our main result.

**Assumption 1.** Take any ball \( B \subset \mathbb{R}^D \) centered on \( \hat{\Theta}(1_T) \) and containing \( \Theta^{(S)} \). We assume the objective \( \text{log} p(x; \Theta, 1_T) - p(\Theta) \) is strongly convex with parameter \( \lambda_{\min} \) on \( B \). Additionally, on \( B \), we assume the derivatives \( g_t(\Theta) := \nabla^{2} \log p(x; \Theta, w)/\partial \Theta \partial w_t \) are Lipschitz continuous with constant \( L_g \) for all \( t \), and the inverse Hessian of the objective is Lipschitz with parameter \( L_{Hinv} \). Finally, on \( B \), take \( \text{log} p(x; \Theta, w_\alpha) \) to be a Lipschitz function of \( \Theta \) with parameter \( L_p \) for all \( w_\alpha \).

We make a few remarks on the restrictiveness of these assumptions. First, while few structured models have objectives that are even convex (e.g., the label switching problem for HMMs guarantees non-convexity), we expect most objectives to be locally convex around an exact minimum \( \hat{\Theta}(1_T); \) Assumption 1 requires that the objective in fact be strongly locally convex. Next, while the Lipschitz assumption on the \( g_t \) may be hard to interpret in general, we note that it takes on a particularly simple form in the setup of Eq. (4), where we have \( g_t = \nabla f_t \). Finally, we note that the condition that the inverse Hessian is Lipschitz is not much of an additional restriction. E.g., if \( \nabla p(\Theta) \) is also Lipschitz continuous, then the entire objective has a Lipschitz gradient, and so its Hessian is bounded. As it is also bounded below by strong convexity, we find that the inverse Hessian is bounded above and below, and thus is Lipschitz continuous. We now state our main result.

**Proposition 2.** The approximation error of \( L_{11}(\Theta^{(S)}) \) satisfies the following bound:

\[ |L_{11}(\Theta^{(S)}) - L_{CV}| \leq C\varepsilon_{\Theta} + \varepsilon_{11}, \]

where \( C := L_p + \frac{L_p L_g}{\lambda_{\min}} + \frac{L_p L_{Hinv}}{|O|} \sum_{\sigma \in O} \left\| \sum_{t \in \sigma} \nabla g_t(\hat{\Theta}(1_T)) \right\|_2 \).

See Appendix J for a proof. Note that, while \( C \) may depend on \( T \) or \( O \), we expect it to approach a constant as \( T \to \infty \) under mild distributional assumptions on \( \|g_t\|_2 \); see Appendix J.

5 Experiments

We demonstrate the effectiveness of our proposed ACV methods on a diverse set of real-world examples where data exhibit temporal and spatial dependence: namely, temporal count modeling, named entity recognition, and spatial modeling of crime data. Additional experiments validating the accuracy and computational benefits afforded by LSCV are available in Appendix K.1, where we explore auto-regressive HMMs for motion capture analysis – with \( N = 124, T \) up to 100, and \( D \) up to 11,712.

**Approximate leave-within-sequence-out CV: Time-varying Poisson processes.** We begin by examining approximate LWCV (Algorithm 1) for maximum a posteriori (MAP) estimation. We consider a time-varying Poisson process model used by \( \text{[Ihler et al. 2006]} \) for detecting events in temporal count data. We analyze loop sensor data collected every five minutes over a span of 25 weeks from a section of a freeway near a baseball stadium in Los Angeles. For this problem, there is one observed sequence (\( N = 1 \)) with \( T = 50,400 \) total observations. There are \( D = 11 \) parameters. Full model details are in Appendix K.1.

To choose the folds in both exact CV and our ACV method, we consider two schemes, both following style (A) in Eq. (4); i.e., we omit observations (but not latents) in the folds. First, we follow the recommendation of \( \text{[Celeux and Durand 2008]} \); namely, we form each fold by selecting \( m \% \) of measurements to omit (i.e., to form \( \sigma \)) uniformly at random and independently across folds. We call this scheme \( \text{i.i.d. LWCV} \). Second, we consider a variant where we omit \( m \% \) of observations in a contiguous block. We call this scheme \( \text{contiguous LWCV} \); see Appendix K.1.

In evaluating the accuracy of our approximation, we focus on a subset of \( T_{sub} = 10,000 \) observations, plotted in the top panel of Fig. 1. The six panels in the lower left of Fig. 1 compare our ACV estimates...
to exact CV. Columns range over left-out percentages $m = 2, 5, 10$ (all on the data subset); rows depict i.i.d. LWCV (upper) and contiguous CV (lower). For each of $|O| = 10$ folds and for each point $x_t$ left out in each fold, we plot a red dot with the exact fold loss $-\log p(x_t | x_{[T]-o}; \hat{\Theta}(w_o))$ as its horizontal coordinate and our approximation $-\log p(x_t | x_{[T]-o}; \hat{\Theta}(w_o))$ as its vertical coordinate. We can see that every point lies close to the dashed black $x = y$ line; that is, the quality of our approximation is uniformly high across the thousands of points in each plot.

In the two lower right panels of Fig. 1 we compare the speed of exact CV to our approximation on two data subsets (size 5,000 and 10,000) and the full data. No reported times include the initial $\hat{\Theta}(1_T)$ computation since $\hat{\Theta}(1_T)$ represents the unavoidable cost of the data analysis itself. I.i.d. LWCV appears in the upper plot, and contiguous LWCV appears in the lower. For our approximation, we use 1,000 folds. Due to the prohibitive cost of exact CV, we run exact CV for 10 folds and multiply by 100 to estimate runtime over 1,000 folds. We see that our approximation confers orders of magnitude in time savings.

Figure 1: Evaluation of approximate LWCV for time-varying Poisson processes. (Top panel) A subset of the count series. (Lower left six panels) Scatter plots comparing exact CV loss (horizontal axis) at each point in each fold (red dots) to our approximation of CV loss (vertical axis). Black dashed line shows perfect agreement. Three columns for percent points left out; two rows for i.i.d. LWCV (upper) and contiguous LWCV (lower). (Lower right two panels) Wall-clock time for exact and approximate CV; same rows as left panels.

Robustness to inexact optimization: Neural conditional random fields. Next, we examine the

Figure 2: Behavior of ACV at different epochs in stochastic optimization for a bilstm-crf. (Four left panels) Scatter plots comparing held out probabilities under CV (horizontal axis) at each point in each fold (red dots) to our approximation of CV (vertical axis). Black dashed line shows perfect agreement. (Right panel) Error in our approximation relative to exact CV and averaged across folds, as a function of log gradient norm in the optimization procedure.
We choose one point per fold in style (B) of LWCV here, for a total of 384 folds. We test our method

We consider 500 LSCV folds with one sentence (i.e., one

We trained the bilstm-crf model on the CoNLL-2003 shared task benchmark [Sang and De Meulder,

−

log

p(x | Θ(Θ(w))) as its horizontal coordinate and our approximation

p(θn | x; Θ(w)) as its vertical coordinate. Note that the LSCV loss has no dependence on other n due to the model independence across n; see Appendix G. Even in early epochs with larger gradient norms, every point lies close to the dashed black x = y line. The rightmost panel in Fig. 2 shows mean absolute approximation error between the exact CV held out probability and our approximation, across all 500 folds and as a function of log gradient norm. Approximation errors as a function of wall clock time are available in Fig. 4. As expected, our approximation has higher quality at better initial fits. Nonetheless, we see that decay in performance away from the exact optimum is gradual.

Beyond chain-structured graphs: Crime statistics in Philadelphia. The models in our experiments above are all chain-structured. Next we consider our approximations to LWCV in a spatial model with more complex dependencies. Balocchi and Jensen [2019], Balocchi et al. [2019] have recently studied spatial models of crime in the city of Philadelphia. We here consider a (simpler) hidden MRF model for exposition: a Poisson mixture with spatial dependencies, detailed in Appendix K.3. Here, there is a single structure observation (N = 1); there are T = 384 census tracts in the city; and there are D = 2 parameters. The data is shown in the upper lefthand panel of Fig. 5.

We choose one point per fold in style (B) of LWCV here, for a total of 384 folds. We test our method across four fixed values of a hyperparameter \( β \) that encourages adjacent tracts to be in the same latent state. For each fold, we plot a red dot comparing the exact fold loss

\[
- \log p(x_t | x_{T-t}; Θ(Θ(w)))
\]

with our approximation

\[
- \log p(x_t | x_{T-t}; Θ(Θ(w)))
\]

The results are in the lower four panels of Fig. 5 where we see uniformly small error across folds in our approximation. In the upper right panel of Fig. 5 we see that our method is orders of magnitude faster than exact CV.

Discussion. In this work, we have demonstrated how to extend approximate cross-validation (ACV) techniques to models with non-trivial dependencies between data points. We have also demonstrated that IJ approximations can retain their usefulness even when the initial data fit is inexact. While our motivation in the latter case was formed by complex models of dependent structures, our results are also new for the classic independence framework of ACV. An interesting remaining challenge for future work is to address other sources of computational expense in structured models. For instance, even after parameter fit, inference can be expensive in very large graphical models; it remains to be seen if reliable and fast approximations can be found for this operation as well.

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CSAIL–MSR Trustworthy AI Initiative, an NSF CAREER Award, an ARO YIP Award, and ONR.
Figure 3: Evaluation of LWCV for loopy Markov random field. (Top left) Census tracts data. (Upper right) Wall-clock time of approximate CV and exact CV. (Lower) Scatter plots comparing CV loss (horizontal axis) at each point in each fold (red dots) to our approximation of CV loss (vertical axis). Black dashed line shows perfect agreement. Plots generated with different values of connectivity $\beta$.

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A Related work: Approximate CV methods

A growing body of recent work has focused on various methods for approximate CV (ACV). As outlined in the introduction, these methods generally take one of two forms. The first is based on taking a single Newton step on the leave-out objective starting from the full data fit, \( \hat{\Theta} \). This approximation was first proposed by Obuchi and Kabashima [2016, 2018] for the special cases of linear and logistic regression and first applied to more general models by Beirami et al. [2017]. While this approximation is generally applicable to any CV scheme (e.g. beyond LOOCV) and any model type (e.g. to structured models), it is only efficiently applicable to LOOCV for GLMs. In particular, approximating each \( \hat{\Theta} o \) requires the computation and inversion of the leave-out objective’s \( D \times D \) Hessian matrix. In the case of LOOCV GLMs, this computation can be performed quickly using standard rank-one matrix updates; however, in more general settings, no such convenience applies.

Various works detail the theoretical properties of the NS approximation. Beirami et al. [2017], Rad and Maleki [2020] provide some of the first bounds on the quality of the NS approximation, but under fairly strict assumptions. Beirami et al. [2017] assume boundedness of both the parameter and data spaces, while Rad and Maleki [2020] require somewhat hard-to-check assumptions about the regularity of each leave-out objective (although they successfully verify their assumptions on a handful of problems). Koh et al. [2019] prove bounds on the accuracy of the NS approximation with fairly standard assumptions (e.g. Lipschitz continuity of higher-order derivatives of the objective function), but restricted to models using \( \ell_2 \) regularization. Wilson et al. [2020] also prove bounds on the accuracy of NS using slightly more complex assumptions but avoiding the assumption of \( \ell_2 \) regularization. More importantly, Wilson et al. [2020] also address the issue of model selection, whereas all previous works had focused on the accuracy of NS for model assessment (i.e. assessing the error of a single, fixed model). In particular, Wilson et al. [2020] give assumptions under which the NS approximation is accurate when used for hyperparameter tuning.

Finally, we note that in its simplest form, the NS approximation requires second differentiability of the model objective. Obuchi and Kabashima [2016, 2018], Rad and Maleki [2020], Beirami et al. [2017], Stephenson and Broderick [2020] propose workarounds specific to models using \( \ell_1 \)-regularization. More generally, Wang et al. [2018] provide a natural extension of the NS approximation to models with either non-differentiable model losses or non-differentiable regularizers.

Again, while these NS methods can be applied to the structured models of interest here, the repeated computation and inversion of Hessian matrices brings their speed into question. To avoid this issue, we instead focus on approximations based on the infinitesimal jackknife (IJ) from the statistics literature [Jaeckel, 1972, Efron, 1981]. The IJ was recently conjectured as a potential approximation to CV by Koh and Liang [2017] and then briefly compared against the NS approximation for this purpose by Beirami et al. [2017]. The IJ was first studied in depth for approximating CV in an empirical and theoretical study by Giordano et al. [2019]. The benefit of the IJ in our application is that for any CV scheme and any (differentiable and i.i.d.) model, the IJ requires only a single matrix inverse to approximate all CV folds. Koh et al. [2019] give further bounds on the accuracy of the IJ approximation for models using \( \ell_2 \) regularization. As in the case for NS, Wilson et al. [2020] give bounds on the accuracy of IJ beyond \( \ell_2 \) regularized models but with slightly more involved assumptions; Wilson et al. [2020] also give bounds on the accuracy of IJ for model selection.

Just as for the NS approximation, the IJ also requires second differentiability of the model objective. Stephenson and Broderick [2020] deal with this issue by noting that the methods of Wang et al. [2018] for applying the NS to non-differentiable objectives can be extended to cover the IJ as well. We note that the use of the IJ for model selection for non-differentiable objectives seems to be more complex than for the NS approximation. In particular, Stephenson and Broderick [2020] Appendix G] show that the IJ approximation can have unexpected and undesirable behavior when

4The methods of Giordano et al. [2019] apply beyond CV to other “reweight and retrain” schemes such as the bootstrap. The methods presented in our paper apply more generally as well, although we do not explore this extension.
used for tuning the regularization parameter for \( \ell_1 \) regularized models. \cite{Wilson2020} resolve this issue by proposing a further modification to the IJ approximation based on proximal operators.

\section{Hidden Markov random fields}

Here we show that HMMs are instances of (hidden) MRFs. Recall that a MRF models the joint distribution,

\[- \log p(x, z; \Theta) = - \sum_{n \in [N]} \log p(x_n, z_n; \Theta) = Z(\Theta) + \sum_{n=1}^{N} \left\{ \sum_{t \in [T]} \psi_t(x_{nt}, z_{nt}; \Theta) \right\} + \left[ \sum_{x \in \mathcal{F}} \phi_x(z_{nt}; \Theta) \right].\]  

\hspace{1cm} (8)

**Hidden Markov models** We recover hidden Markov models as described in Section 2 by setting \( \psi_t(x_{nt}, z_{nt}; \Theta) = \log F(x_{nt} \mid z_{nt}) \), setting \( \mathcal{F} \) to the set of all unary and pairwise indices, and defining \( \phi_{t,t-1}(z_{nt}, z_{nt-1}; \Theta) = \log \text{Cat}(z_{nt} \mid A_{z_{nt-1}}) \), and \( \phi_1(z_{n1}) = \log \text{Cat}(z_{n1} \mid \pi) \), and \( \phi_t(z_{nt}) = 0 \), for \( t \in [T] - 1 \). The log normalization constant is \( Z(\Theta) = 0 \).

\section{Leave structure out cross-validation (LSCV)}

**Algorithm 2 Structured approximate LSCV**

\begin{align*}
\text{Require: } & \Theta_1, x, \mathcal{O} \\
1: & \text{Marginalize over } z_n: \log p(x_n; \Theta) = \text{MARG}(x_n; \Theta), \forall n \in [N] \\
2: & \text{Compute } \log p(x; \Theta, w) = \sum_n w_n \log p(x_n; \Theta) \\
3: & \text{Compute } H = \left. \frac{\partial^2 \log p(x; \Theta, w) + \log p(\Theta)}{\partial \Theta \partial \Theta} \right|_{\Theta=\Theta_1, w=1_N} \\
4: & \text{Compute } J = (J_{dn}) := \left. \frac{\partial^2 \log p(x; \Theta, w) + \log p(\Theta)}{\partial \Theta \partial w_n} \right|_{\Theta=\Theta_1, w=1_N} \\
5: & \text{for } o \in \mathcal{O}, \hat{\Theta}_{\text{ACV}}(\Theta_1, x, o) := \Theta_1 + \sum_{n \in o} H^{-1} J_n \quad \# J_n \text{ is } n\text{th column of } J \\
6: & \text{return } \{\hat{\Theta}_{\text{ACV}}(\Theta_1, x, o)\}_{o \in \mathcal{O}}
\end{align*}

\section{Efficient weighted marginalization (WEIGHTEDMARG) for chain-structured MRFs}

For chain-structured pairwise MRFs with discrete structure, we can use a dynamic program to efficiently marginalize out the structure. Assume that \( z_t \forall t \in [T] \) can take one of \( K \) values. Define \( \alpha_{t1} = \exp\{w_1 \psi_1(x_1, z_1 = k)\} \), and then compute \( \alpha_{t,k} \) recursively:

\[ \alpha_{t,k} = \sum_{\ell=1}^{K} \alpha_{t-1,\ell} \exp \left[ w_t \psi_t(x_t, z_t = k; \Theta) + \phi_{t,t-1}(z_t = k, z_{t-1} = \ell) \right], \]  

\hspace{1cm} (9)

if using weighting scheme (A) from Equation Eq. (4) or,

\[ \alpha_{t,k} = \sum_{\ell=1}^{K} \alpha_{t-1,\ell} \exp \left[ w_t \psi_t(x_t, z_t = k; \Theta) + w_t w_{t-1} \phi_{t,t-1}(z_t = k, z_{t-1} = \ell) \right], \]  

\hspace{1cm} (10)

if using weighting scheme (B) from Equation Eq. (5). Then, for either (A) or (B), we have \( p(x; \Theta, w) = \sum_{k=1}^{K} \alpha_{T,k} \). When \( w = 1_T \) we recover the empirical risk minimization solution. As is the case for non-weighted models, this recursion implies that \( p(x; \Theta, w) \) is computable in \( O(TK^2Q) \) time instead of the usual \( O(T^2K) \) time required by brute-force summation (recall \( Q \) is the time required to evaluate one local potential). Likewise, we can also compute the derivatives needed by Algorithm 1 in \( O(TK^2Q) \) time either by manual implementation or automatic differentiation tools \cite{Bartholomew-Biggs2000}.
E  Equivalence of weighting (A) and (B) for leave-future-out for chain-structured graphs

As noted in the main text, weighting schemes (A) and (B) are equivalent when the graph is chain structured. Formally,

**Proposition 3.** Consider a chain-structured pairwise MRF with ordered indices $t$ on the chain (such as an HMM). Weighting styles (A) and (B) above are equivalent for leave-future-out CV. That is, choose $o = \{T', \ldots, T\}$ for some $T' \in [T-1]$ (i.e., indices that are in the “future” when interpreted as time). Then set $\forall t \in o, w_t = 0$ and $\forall t \in [T] - o, w_t = 1$.

This result does not hold generally beyond chain-structured graphical models – consider a four-node “ring” graph in which node $t$ is connected to nodes $t-1$ and $t+1$ (mod 4) for $t = 0, \ldots, 3$. Weighting scheme (B) produces a distribution that is chain-structured over three nodes, whereas (A) produces a distribution without such conditional independence properties. We now prove Proposition 3.

**Proof.** Recall that for a chain structured graph, we can write:

$$p(x, z) = p(x \mid z) p(z_1) \prod_{t=2}^{T} p(z_t \mid z_{t-1}).$$

Let $o = \{T', T' + 1, \ldots, T\}$ for some $T' < T$; that is, we are interested in dropping out time steps $T', \ldots, T$. For weighting scheme (A) (Eq. (4)), we drop out only the observations, obtaining:

$$p_A(x, z; w_o) = \left( \prod_{t=1}^{T'-1} p(x_t \mid z_t) \right) p(z_1) \prod_{t=2}^{T} p(z_t \mid z_{t-1}).$$

When we sum out all $z$ to compute the marginal $p_A(x; w_o)$, we can first sum over $z_T, \ldots, z_{T'}$. As $\sum_z p(z_t \mid z_{t-1}) = 1$ for any value of $z_{T-1}$, we obtain:

$$p_A(x; w_o) = \sum_{z_1, \ldots, z_{T'-1}} \left( \prod_{t=1}^{T'-1} p(x_t \mid z_t) \right) p(z_1) \prod_{t=2}^{T'-1} p(z_t \mid z_{t-1}),$$

which is exactly the formula for $p_B(x; w_o)$, the marginal likelihood from following weighting scheme (B) (Eq. (5)), in which we drop out both the $x_t$ and $z_t$ for $t \notin o$. \qed

F  Conditional random fields

Conditional random fields assume that the labels $z$ are observed and model the conditional distribution $p(z \mid x; \Theta)$. While more general dependencies between $x$ and $z$ are possible a commonly used variant [Ma and Hovy 2016, Lample et al. 2016] captures the conditional distribution of the joint defined in Equation Eq. (8). Note,

$$\log p(z_n \mid x_n; \Theta) = \log p(x_n, z_n; \Theta) - \log p(x_n; \Theta)$$

$$= -Z(\Theta) + \sum_{t \in [T]} \psi_t(x_{nt}, z_{nt}; \Theta) + \sum_{e \in F} \phi_e(z_{ne}; \Theta)$$

$$+ Z(\Theta) - \int_{z_n} \sum_{t \in [T]} \psi_t(x_{nt}, z_{nt}; \Theta) + \sum_{e \in F} \phi_e(z_{ne}; \Theta) dz_n$$  \hspace{1cm} (11)

Defining, $Z(x_n; \Theta) = -\int_{z_n} \sum_{t \in [T]} \psi_t(x_{nt}, z_{nt}; \Theta) + \sum_{e \in F} \phi_e(z_{ne}; \Theta) dz_n$, then gives us the following conditional distribution,

$$-\log p(z \mid x; \Theta) = \sum_{n=1}^{N} \left\{ Z(x_n; \Theta) + \sum_{t \in [T]} \psi_t(x_{nt}, z_{nt}; \Theta) + \sum_{e \in F} \phi_e(z_{ne}; \Theta) \right\}.$$  \hspace{1cm} (12)

Note that $Z(x_n; \Theta)$ is an observation specific negative normalization constant.
CV for conditional random fields

Analogously to the MRF case, we have two variants for CRFs — LSCV and LWCV. While LSCV is frequently used in practice, for example, [DeCaprio et al., 2007], we are unaware of instances of LWCV in the literature. Thus, while we derive approximations to both CV schemes, our CRF-based experiments in Section 5 only use LSCV.

G.1 LSCV for CRFs

Leave structure out CV is analogous to the MRF case and is detailed in Algorithm 3, where

$$\log \hat{p}(z_n, x_n; \Theta) := \sum_{t \in [T]} \psi_t(x_{nt}, z_{nt}; \Theta) + \sum_{c \in F} \phi_c(z_{nc}; \Theta).$$

Since all input, label pairs \(\{x_n, z_n\}\) are independent, \(\log p(z \mid x; \Theta, w)\) is just a weighted sum across \(n\) and the losses \(-\log p(z_n \mid x_n; \hat{\Theta}(w_{\{n\}}))\) and \(-\log p(z_n \mid x_n; \hat{\Theta}(w_{\{n\}}))\) do not depend on \([N] - n\).

Algorithm 3 Structured approximate cross-validation (LSCV) for CRFs

Require: \(\Theta_1, x, z, O\)

1: Compute \(Z(x_n; \Theta) = -\text{MARG}(x_n; \Theta), \forall n \in [N]\)
2: Compute \(\log p(z \mid x; \Theta, w) = \sum_n w_n [Z(x_n; \Theta) + \log \hat{p}(z_n, x_n; \Theta)]\)
3: Compute \(H = \left(\frac{\partial^2 \log p(z \mid x; \Theta, w) + \log p(\Theta)}{\partial \Theta \partial \Theta} \right)_{\Theta=\Theta_1, w=1_N}\)
4: Compute matrix \(J := (J_{dn}) = \left(\frac{\partial^2 \log p(z \mid x; \Theta, w) + \log p(\Theta)}{\partial \Theta \partial w_n} \right)_{\Theta=\Theta_1, w=1_N}\)
5: for \(o \in O\), \(\hat{\Theta}_{ACV}(\Theta_1, x, z, o) := \Theta_1 + \sum_{n \in o} H^{-1} J_n\) # \(J_n\) is \(n\)th column of \(J\)
6: return \(\{\hat{\Theta}_{ACV}(\Theta_1, x, z, o)\}_{o \in O}\)

G.2 LWCV for CRFs

Leave within structure out for CRFs again comes with a choice of weighting scheme. Given a single input, label pair \(x, z\), the \(z_t\) are the outputs at location \(t\), and the \(x_t\) are the corresponding inputs. A form of CV arises when we drop the outputs \(z_t\), for \(t \in o\). This gives us weighting scheme (C),

$$\hat{\Theta}(w) = \arg\min_{\Theta} Z(\Theta, w, x)$$

$$+ \left[ \sum_{t \in [T]} w_t \psi_t(x_t, z_t; \Theta) + (1 - w_t) \int_{z_t} \psi_t(x_t, z_t; \Theta) \, dz_t \right]$$

$$+ \left[ w_t \sum_{c \in F} \phi_c(z_c; \Theta) + (1 - w_t) \int_{z_t} \sum_{c \in F} \phi_c(z_c; \Theta) \, dz_t \right] - \log p(\Theta).$$

For linear chain structured CRFs with discrete outputs \(z\) a variant of the forward algorithm can be used to efficiently compute \(Z(\Theta, w, x)\) as well as the marginalizations over \(\{z_t \mid t \in o\}\) required by Eq. (13). See Bellare and McCallum [2007], Tsiboi et al. [2008] for details. Algorithm 4 summarizes the steps involved.

H Computational cost of one Newton-step-based ACV

Recall that we define \(M\) to be the cost of one marginalization over the latent structure \(z\) and noted above that the cost of computing the Hessian via automatic differentiation is \(O(M)\). For the Newton step approximation, recall that we need to compute a different Hessian for each fold \(o\). While this can be avoided using rank-one update rules in the case of leave-one-out CV for generalized linear models, this is not the case for the CV schemes and models considered here. Thus, to use the Newton step approximation here, we require \(O(M|O|)\) time to compute all needed Hessians. Compared to the \(O(M)\) time spent computing Hessians by our algorithms, the Newton step is significantly more expensive. For this reason, we do not consider Newton step based approximations here.
We prove here a slightly more general version of Proposition 2 that covers both LWCV and LSCV, as well as arbitrary loss functions $\ell$. To encompass both in the same framework, let $w_n \in \mathbb{R}^T$ be weight vectors for each structured object $n = 1, \ldots, N$. Our weighted objective will be:

$$\hat{\Theta}(w) = \arg\min_{\Theta \in \mathbb{R}^D} \sum_{n=1}^{N} \log p(D_n; \Theta, w_n) + p(\Theta),$$

**Algorithm 4** Approximate leave-within-structure-out cross-validation for CRFs

**Require**: $\Theta_1, x, z, O$

1. Compute *unweighted* marginalization over $z$, $Z(x; \Theta) = -\text{MARG}(x_n; \Theta)$, $\forall n \in [N]$.
2. Compute *weighted* marginalization over $z$: $Z(x; \Theta, w) = \text{WEIGHTEDMARG}(x, \Theta, w)$.
3. Compute $\log p(z | x; \Theta) = Z(x; \Theta, w) + Z(x_n; \Theta)$
4. Compute $H = \frac{\partial^2 \log p(z | x, \Theta, w)}{\partial \Theta \partial \Theta^T} + \log p(\Theta)_{\Theta=\hat{\Theta}_1, w=1_T}$
5. Compute matrix $J := (J_\Theta) = \left( \frac{\partial \log p(x, \Theta, w) + \log p(\Theta)}{\partial \Theta} \right)_{\Theta=\hat{\Theta}_1, w=1_T}$
6. for $o \in O$. do: $\hat{\Theta}_{\text{ACV}}(\Theta_1, x, o) := \Theta_1 + \sum_{t \in o} H^{-1} J_t$ # $J_t$ is $t$th column of $J$
7. return $\{\hat{\Theta}_{\text{ACV}}(\Theta_1, x, o)\}_{o \in O}$

I Derivation of IJ approximations

In all cases considered here (i.e., the “exchangeable” leave-one-out CV considered by previous work or the more structured variants for chain-structured or general graph structured models) can be derived similarly. In particular, once we have derived the relevant weighted optimization problem for each case, the derivation of the IJ approximation is the same. Let the relevant weighted optimization problem be defined for $w \in \mathbb{R}^T$:

$$\hat{\Theta}(w) := \arg\min_{\Theta \in \mathbb{R}^D} F(\Theta, w),$$

where $F$ is some objective function with $F(\cdot, 1_T)$ corresponding to the “full-data” fit (i.e., without leaving out any data). We now follow the derivation of the IJ in Giordano et al. [2019]. The condition that $\Theta(1_T)$ is an exact optimum is:

$$\left. \frac{\partial F}{\partial \Theta} \right|_{\hat{\Theta}(1_T), 1_T} = 0.$$

If we take a derivative with respect to $w_t$:

$$\left. \frac{\partial^2 F}{\partial \Theta \partial \Theta^T} \right|_{\hat{\Theta}(1_T), 1_T} d\Theta \
\left. \frac{\partial^2 F}{\partial \Theta \partial w_t} \right|_{\hat{\Theta}(1_T), 1_T} d\Theta + \left. \frac{\partial^2 F}{\partial \Theta \partial w_t} \right|_{\hat{\Theta}(1_T), 1_T} d\Theta \
\left. \frac{\partial^2 F}{\partial \Theta \partial w_t} \right|_{\hat{\Theta}(1_T), 1_T} = 0.$$

Noting that $d\Theta/d\Theta = 1$ and solving for $d\Theta/dw_t$:

$$\left. \frac{d\Theta}{dw_t} \right|_{\hat{\Theta}(1_T), 1_T} = - \left( \left. \frac{\partial^2 F}{\partial \Theta \partial \Theta^T} \right|_{\hat{\Theta}(1_T), 1_T} \right)^{-1} \left. \frac{\partial^2 F}{\partial \Theta \partial w_t} \right|_{\hat{\Theta}(1_T), 1_T}$$

(14)

Thus we can form a first order Taylor series of $\hat{\Theta}(w)$ in $w$ around $w = 1_N$ to approximate:

$$\hat{\Theta}_{1J}(w) \approx \hat{\Theta}(1_T) - \sum_{t=1}^{T} \left( \left. \frac{\partial^2 F}{\partial \Theta \partial \Theta^T} \right|_{\hat{\Theta}(1_T), 1_T} \right)^{-1} \left. \frac{\partial^2 F}{\partial \Theta \partial w_t} \right|_{\hat{\Theta}(1_T), 1_T} (1 - w_t).$$

Specializing this last equation to the various $F$ and weight vectors $w$ of interest derives each of our ACV algorithms.

J Inexact optimization

We prove here a slightly more general version of Proposition 2 that covers both LWCV and LSCV, as well as arbitrary loss functions $\ell$. To encompass both in the same framework, let $w_n \in \mathbb{R}^T$ be weight vectors for each structured object $n = 1, \ldots, N$. Our weighted objective will be:

$$\hat{\Theta}(w) = \arg\min_{\Theta \in \mathbb{R}^D} \sum_{n=1}^{N} \log p(D_n; \Theta, w_n) + p(\Theta),$$
where \( D = \{ D_1, \ldots, D_N \} \) denotes the collection of all observed structures; i.e., each \( D_n \) may be a sequence of observations \( x_n \) for a HMM or observed outputs and inputs \( x_n, z_n \) for a CRF. Let \( \hat{\Theta}(1_{NT}) \) be the solution to this problem with \( w_{nt} = 1 \) for all \( n \) and \( t \). We assume that we are interested in estimating the exact out-of-sample loss for some generic loss \( \ell \) by using exact CV, \( \mathcal{L}_{CV} := \frac{1}{|O|} \sum_o \ell(D_o, \mathcal{D}_o, \hat{\Theta}(w_o)) \); e.g., we may have \( \ell(D_o, \mathcal{D}_o, \hat{\Theta}(w_o)) = -\log p(x_o \mid x_{\mathcal{D}_o}, \hat{\Theta}(w_o)) \) in the case of a HMM with \( N = 1 \). Notice here that \( o \in [N] \times [T] \) indexes arbitrarily across structures. We can now state a modified version of Assumption 1.

Assumption 2. Let \( B \subset \mathbb{R}^D \) be a ball centered on \( \hat{\Theta}(1_{NT}) \) and containing \( \Theta(S) \). Then the objective \( \sum_n \log p(x_n; \Theta, 1_T) + p(\Theta) \) is strongly convex with parameter \( \lambda_{\text{min}} \) on \( B \). Additionally, on \( B \), the derivatives \( g_{nt}(\Theta) := \frac{\partial^2}{\partial \Theta \partial w_n} \log p(x_n; \Theta, w_n) \) are Lipschitz continuous with constant \( L_{\ell} \) for all \( n, t \) and the inverse Hessian of the objective is Lipschitz with parameter \( L_{H_{\text{inv}}} \). Finally, on \( B \), \( \ell(D_o, \mathcal{D}_o, \Theta) \) is a Lipschitz function of \( \Theta \) with parameter \( L_{\ell} \) for all \( o \).

We now prove our more general version Proposition 2.

Proposition 4. Take Assumption 2. Then the approximation error of \( \mathcal{L}_{11}(\Theta(S)) \) is bounded by:

\[
|\mathcal{L}_{11}(\Theta(S)) - \mathcal{L}_{CV}| \leq C \varepsilon_{\Theta} + \varepsilon_{11},
\]

where \( C \) is given by

\[
\left( L_{\ell} + \frac{L_{\ell} L_g}{\lambda_{\text{min}}} + \frac{L_{\ell} L_{H_{\text{inv}}}}{|O|} \sum_o \left\| \sum_{t \in o} g_{nt}(\hat{\Theta}(1_{NT})) \right\|_2 \right).
\]

Proof. By the triangle inequality:

\[
|\mathcal{L}_{11}(\Theta(S)) - \mathcal{L}_{CV}| \leq
\]

\[
|\mathcal{L}_{11}(\Theta(S)) - \mathcal{L}_{11}(\hat{\Theta}(1_{NT}))| + |\mathcal{L}_{11}(\hat{\Theta}(1_{NT})) - \mathcal{L}_{CV}|.
\]

The second term is just the constant \( \varepsilon_{11} \). Now we just need to bound the first term using our Lipschitz assumptions. We have, by the triangle inequality

\[
|\mathcal{L}_{11}(\hat{\Theta}(1_{NT})) - \mathcal{L}_{11}(\Theta(S))| \leq \frac{1}{|O|} \sum_o \left| \ell \left( D_o, \mathcal{D}_o, \hat{\Theta}(1_{NT}) + H^{-1}(\hat{\Theta}(1_{NT})) \sum_{t \in o} g_{nt}(\hat{\Theta}(1_{NT})) \right) \right. \\
- \left. \ell \left( D_o, \mathcal{D}_o, \Theta(S) + H^{-1}(\Theta(S)) \sum_{t \in o} g_{nt}(\Theta(S)) \right) \right|.
\]

Continuing to apply the triangle inequality and our Lipschitz assumptions:

\[
\leq \frac{L_{\ell}}{|O|} \sum_o \left( \left\| \hat{\Theta}(1_{NT}) - \hat{\Theta}(S) \right\|_2 + \left\| H^{-1}(\hat{\Theta}(1_{NT})) \sum_{t \in o} g_{nt}(\hat{\Theta}(1_{NT})) - H^{-1}(\Theta(S)) \sum_{t \in o} g_{nt}(\Theta(S)) \right\|_2 \right)
\]

\[
\leq L_{\ell} \varepsilon_{\Theta} + \frac{L_{\ell}}{|O|} \sum_o \left\| \sum_{t \in o} \left( H^{-1}(\hat{\Theta}(1_{NT})) \sum_{t \in o} g_{nt}(\hat{\Theta}(1_{NT})) - H^{-1}(\Theta(S)) \sum_{t \in o} g_{nt}(\Theta(S)) \right) \right\|_2
\]

\[
\leq L_{\ell} \varepsilon_{\Theta} + \frac{L_{\ell} L_g}{\lambda_{\text{min}}} \sum_o \left\| \sum_{t \in o} g_{nt}(\hat{\Theta}(1_{NT})) \right\|_2 \varepsilon_{\Theta}.
\]

Defining the term in the parenthesis as \( C \) finishes the proof.
As noted after the statement of Proposition 2 in the main text, \((1/|O|) \sum_{o \in O} \sum_{t \in o} g_{nt}(\hat{\Theta}(1_{NT}))\) may depend on \(T\), \(N\) or \(O\), but we expect it to converge to a constant given reasonable distributional assumptions on the data. To build intuition, we consider the case of leave-one-out CV for generalized linear models, where we observe a dataset of size \(N > 1\) and have \(T = 1\). In particular, we have \(\log(x_n, y_n; \Theta) = f(x_n^T \Theta, y_n)\), where \(x_n \in \mathbb{R}^D\) are the covariates and \(y_n \in \mathbb{R}\) are the responses. In this case, \(g_{nt} = D_n^{(1)} x_n\), where \(D_n^{(1)} = df(z)/dz\) \(\bigg|_{z=x_n^T \hat{\Theta}(1_T)}\). Then, given reasonable distributional assumptions on the covariates and some sort of control over the derivatives \(D_n^{(1)}\), we might suspect that \((1/N) \sum_{n \in |O|} D_n^{(1)} \|x_n\|_2\) will converge to a constant. As an example, we consider logistic regression with sub-Gaussian data, for which we can actually prove high-probability bounds on this sum.

**Definition 1.** ([Vershynin, 2018]) For \(c_x > 0\), a random variable \(V\) is \(c_x\)-sub-Gaussian if

\[
\mathbb{E} \left[ \exp \left( V^2 / c_x^2 \right) \right] \leq 2.
\]

**Proposition 5.** For logistic regression, assume that the components of the covariates \(x_n d\) are i.i.d. from a zero-mean \(c_x\)-sub-Gaussian distribution for \(d = 1, \ldots, D\). Then we have that, for any \(t \geq 0\):

\[
\Pr \left[ \left\| \nabla f(\hat{\Theta}(1_T), x_n) \right\|_2 - \sqrt{D} \right| \geq t \right] \leq \exp \left[ -C \frac{N^2 t^2}{c_x^2} \right], \tag{16}
\]

where \(C > 0\) is some global constant, independent of \(N, D,\) and \(c_x\).

**Proof.** First, we can use the fact that \(\|\nabla f(\hat{\Theta}(1_T), x_n)\|_2 \leq \|x_n\|_2\), as for logistic regression, \(|D_n^{(1)}| \leq 1\). Next, we can use the fact that \(\|x_n\|_2 - \sqrt{D}\) is a zero-mean sub-Gaussian random variable by Theorem 3.1.1 of [Vershynin, 2018]. We can then apply Hoeffding’s inequality ([Vershynin, 2018], Theorem 2.6.3) to complete the proof.

### K Experimental details

We provide further experimental details in this section.

#### K.1 Time varying Poisson processes

We briefly summarize the time-varying Poisson process model from [Ihler et al., 2006] here. Our data is a time series of loop sensor data collected every five minutes over a span of 25 weeks from a section of a freeway near a baseball stadium in Los Angeles. In all, there are 50,400 measurements of the number of cars on that span of the freeway. Ihler et al. analyze the resulting time series of counts \(x\) to detect the presence or absence of an event at the stadium. Following their model, we use a background Poisson process with a time varying rate parameter \(\lambda_t\) to model non-event counts, \(x_{b_t} \sim \text{Poisson}(\lambda_t)\).

To model the daily variation apparent in the data, we define \(\lambda_t = \lambda_0 \delta_{d_t}\), where \(d_t\) takes one of seven values, each corresponding to one day of the week and \(\delta_{\{1/7, \ldots, 6/7\}} \sim \text{Dir}(1, \ldots, 1)\). We use binary latent variables \(z_t\) indicate the presence or absence of an event and assume a first order Markovian dependence, \(z_t | z_{t-1} \sim A z_{t-1}\). Next, \(z_t = 0\) indicates a non-event at time step \(t\) and the observed counts are generated as \(x_t = x_{b_t}\). An event at time step \(t\) corresponds to \(z_t = 1\) and \(x_t = x_{b_t} + x_{e_t}\), and \(x_{e_t} \sim \text{NegBinomial}(x_{e_t} | a, b/(1 + b))\), where \(x_{e_t}\) are unobserved excess counts resulting from the event. We place Gamma priors on \(\lambda_0, a\) and Beta priors on \(A_{00}\) and \(A_{11}\), and learn the MAP estimates of the parameters \(\Theta = \{\lambda_0, \delta_1, \ldots, \delta_7, a, b, A\}\) while marginalizing \(x_{e_t}\) and \(z_1, \ldots, z_T\). We refer the interested reader to [Ihler et al., 2006] for further details about the model and data.

**Contiguous LWCV.** In contiguous LWCV we leave out contiguous blocks from a time series. To drop \(m\%\) of the data, we sample an index \(t\) uniformly at random from \(\lfloor mT/100 \rfloor + 1, \ldots, T\) and set \(o = \{t - \lfloor mT/100 \rfloor, \ldots, t\}\).

#### K.2 Neural CRF

We employed a bi-directional LSTM model with a CRF output layer. We used a concatenation of a 300 dimensional Glove word embeddings ([Pennington et al., 2014]) and a character CNN ([Ma and...]}
Hovy [2016] based character representation. We employed variational dropout with a dropout rate of 0.25. The architecture is detailed below.

```
LSTMCRFVD(
    (dropout): Dropout(p=0.25, inplace=False)
    (char_feats_layer): CharCNN(
        (embedding): Embedding(96, 50, padding_idx=0)
        (embedding_dropout): Dropout(p=0.25, inplace=False)
    )
    (cnn): Conv1d(50, 30, kernel_size=(3,), stride=(1,), padding=(2,))
    (word_embedding): Embedding(2196016, 300)
    (rnn): StackedBidirectionalLstm(
        (forward_layer_0): AugmentedLstm(
            (input_linearity): Linear(in_features=330, out_features=200, bias=False)
            (state_linearity): Linear(in_features=50, out_features=200, bias=True)
        )
        (backward_layer_0): AugmentedLstm(
            (input_linearity): Linear(in_features=330, out_features=200, bias=False)
            (state_linearity): Linear(in_features=50, out_features=200, bias=True)
        )
        (forward_layer_1): AugmentedLstm(
            (input_linearity): Linear(in_features=100, out_features=200, bias=False)
            (state_linearity): Linear(in_features=50, out_features=200, bias=True)
        )
        (backward_layer_1): AugmentedLstm(
            (input_linearity): Linear(in_features=100, out_features=200, bias=False)
            (state_linearity): Linear(in_features=50, out_features=200, bias=True)
        )
        (layer_dropout): InputVariationalDropout(p=0.25, inplace=False)
    )
    (rnn_to_crf): Linear(in_features=100, out_features=9, bias=True)
    (crf): ConditionalRandomField()
)
```

Training. We used Adam for optimization. Following the recommendation of Reimers and Gurevych [2017] we used mini-batches of size 31. We employed early stopping by monitoring the loss on the validation set. Freezing all but the CRF layers we further fine-tuned only the CRF layer for an additional 60 epochs. In Fig. 4 we plot the mean absolute approximation error in the held out probability under exact CV and our approximation across all 500 folds as a function of (wall clock) time taken by the optimization procedure.

K.3 Philadelphia crime experiment

Our crime data comes from opendataphilly.org where the Philadelphia Police Department publicly releases the time, type, and location of every reported time. For each census tract, we have a latent label $z_t \in \{-1, 1\}$, and model the number of reported crimes $x_t$ with a simple Poisson mixture model: $x_t | z_t \sim \text{Poisson}(\lambda_{z_t})$ where $\lambda_{-1}, \lambda_1 > 0$ are the unknown mean levels of crime in low- and high-crime areas, respectively. Since we might expect adjacent census tracts to be in the same latent state, we model the $z_t$’s with an MRF so that

$$\log p(x, z; \Theta) = \sum_t [-\lambda_{z_t} + x_t \log \lambda_{z_t} - \log(x_t)] + \beta \sum_t \sum_{t' \in \Gamma(t)} 1\{z_t = z_{t'}\} - \log Z(\beta)$$

where $\Theta = \{\lambda_{-1}, \lambda_1\}, \Gamma(t)$ is the collection of census tracts that are spatially adjacent to census tract $t$ and $\log Z(\beta)$ is the log normalizer for the latent field $p(z)$. The potential $1\{z_t = z_{t'}\}$ expresses prior belief that adjacent census tracts should be in the same latent class. The connection strength $\beta$ is treated as a hyper-parameter. For each $\beta$ fixed, $\Theta$ is estimated using expectation maximization [Dempster et al., 1977] on $\sum_z \log p(x, z; \Theta)$. M-step computation is analytical, given the posteriors...
Figure 4: Error in our approximation relative to exact CV averaged across folds, as a function of wall clock time.

$p(z_t | x_t; \Theta)$. Exact E-step computation is reasonably efficient through smart variable elimination [Koller and Friedman, 2009, Chapter 9]: the number of states is small and common heuristics to find good elimination orderings, such as MinFill, worked well. This efficient variable elimination order is also used to implement the WEIGHTEDMARG routine of [1].

L Additional experiments

We present additional experimental validation in support of the ACV methods in this section.

L.1 Motion capture analysis

Data. We analyze motion capture recordings from the CMU MoCap database (http://mocap.cs.cmu.edu), which consists of several recordings of subjects performing a shared set of activities. We focus on the 124 sequences from the “Physical activities and Sports” category that has been previously been studied [Fox et al., 2009, Hughes et al., 2012, Fox et al., 2014] in the context of unsupervised discovery of shared activities from the observed sequences. At each time step we retain twelve measurements deemed informative for describing the activities of interest, as recommended by Fox et al.. Auto-regressive hidden Markov models have been shown effective for this task, motivating their use in this section.

Accurate LSCV—auto-regressive HMMs We confirm here that ACV is accurate and computationally efficient for structured models in the case studied by previous work: LSCV with exact model fits. We present comparisons between embarrassingly parallel exact CV and LSCV with parallelized Hessian computation (“Approx. Parallel”, i.e., we parallelize the Hessian computation over different structures $n$), alleviating the primary computational bottleneck for ACV. We model the collection of MoCAP sequences via a $K$-state HMM with an order-$p$ auto-regressive (AR(p)) observation model. We also consider variants where each state’s auto-regressive model is parameterized via a neural network. Figure 5 visualizes a MoCAP sequence where we have retained only the 12 relevant dimensions. For this experiment, we retain up to 100 (= $T$) measurements per sequence. We employ the following auto-regressive observation model,

$$p(x_{nt} | x_{nt-1}, \ldots, x_{nt-p}, z_{nt}) = \mathcal{N}(x_{nt} | \sum_{m=1}^{p} B^1_{z_{nt}} x_{nt-m} + b^1_{z_{nt}}, \sigma^2 I),$$

$$B_k \sim \text{Matrix-Norm}(I, I, I), \quad b_k \sim \mathcal{N}(0, I), \quad \forall k \in \{1, \ldots, K\},$$

where $p$ is the order of the auto-regression. Neural auto-regressive observation models are defined as,

$$p(x_{nt} | x_{nt-1}, \ldots, x_{nt-p}, z_{nt}) = \mathcal{N}(x_{nt} | B^0_{z_{nt}} h(\sum_{m=1}^{p} B^0_{z_{nt}} x_{nt-m} + b^0_{z_{nt}}) + b^1_{z_{nt}}, \sigma^2 I),$$

$$\theta_k \sim \mathcal{N}(0, \lambda I), \quad \forall k \in \{1, \ldots, K\},$$

\(20\)
Figure 5: Motion capture analysis through auto-regressive HMMs. (Top) A twelve dimensional MoCap sequence that serves as the observed data and the number of parameters $D$ for different models under consideration. The high dimensionality of the models make alternate ACV methods based on a single Newton step infeasible. (Middle) Scatter plots comparing leave one out loss, where x-axis is $-\ln p(x_n \mid \Theta(w_{(n)}))$ and y-axis is $-\ln p(x_n \mid \hat{\Theta}_I J(w_{(n)}))$ for different auto-regressive orders under exact and IJ approximated leave one out cross validation. Points along the diagonal indicate accurate IJ approximations. (Bottom) Timing and held out negative log probability across different models. For IJ and Exact the error bars represent two jackknife standard error. The IJ approximations are significantly faster but closely approximate exact leave one out loss across models and track well with test loss computed on the held out $20\%$ of the dataset.

where $\theta_k = \{B_0^k, b_0^k, B_1^k, b_1^k\}$, and $h$ denotes a tanh non-linearity, and $B_{k}^{0}, B_{k}^{1} \in \mathbb{R}^{12 \times 12}$ and $b_{k}^{0}, b_{k}^{1} \in \mathbb{R}$, i.e., a $12$-$12$-$12$ fully connected network.

While past work has explored AR(0) and AR(1) observation models, a thorough exploration of the effect of $p$ has been lacking. ACV provides an effective tool for exploring such questions accurately and inexpensively. We split the sequences into a $80/20\%$ train and test split and perform LSCV on the training data ($N = 100$) to compare AR(p) models with $p$ ranging from zero through five and the neural variant with $p = 1$ (NAR(1)), in terms of how well they describe the left out sequence. Following Fox et al., we fix $K = 16$. Figure 5 summarizes our results. First, we see that the ACV loss is quite close to the exact CV loss and that both track well with the held-out test loss. Furthermore, consistent with previous studies, we find that using an AR(1) observation model is significantly better than using an AR(0) or higher-order AR model. Interestingly, the out-of-sample loss for the AR(1) model is comparable to neural variant, NAR(1).

In terms of computation, the ACV is significantly faster than exact CV. In fact, for the higher order auto-regressive likelihoods and the neural variant, exact CV was too expensive to perform. Instead, we report estimated time for running such experiments by multiplying the average time taken to run three folds of LSCV with the number of training instances. For AR(0) and AR(1) we compare against exact CV implemented via publicly available optimized Expectation Maximization code [Hughes and Sudderth, 2014]. The higher order AR and the NAR(1) model, were fit by BFGS as implemented in SCIPY.OPTIMIZE.MINIMIZE. We find that computing the embarrassingly parallel version provides significant speedups over their serial counterparts.
Within sequence leave out experiments. We took the longest MoCAP sequence containing 1484 measurements and fit a five state HMM with Gaussian emissions. We find that even for the MoCAP data IJ approximations to i.i.d. LWCV is very accurate. As the contiguous LWCV involves making larger scale changes to the sequence, for instance at 10% we end up dropping chunks of 140 time steps from the sequences, resulting in larger changes to the parameters, IJ approximations are relatively less accurate.

Figure 6: Within sequence leave out experiments. We took the longest MoCAP sequence containing 1484 measurements and fit a five state HMM with Gaussian emissions. We find that even for the MoCAP data IJ approximations to i.i.d. LWCV is very accurate. As the contiguous LWCV involves making larger scale changes to the sequence, for instance at 10% we end up dropping chunks of 140 time steps from the sequences, resulting in larger changes to the parameters, IJ approximations are relatively less accurate.

(Top) Scatter plots comparing i.i.d LWCV loss $-\ln p(x_t | x_{[T]-o}; \Theta(\omega))$ (horizontal axis) with $-\ln p(x_t | x_{[T]-o}; \hat{\Theta}_{IJ}(\omega))$ (vertical axis), for each point $x_t$ left out in each fold, computed under exact CV for different omission rates $m\% = 2\%, 5\%, 10\%$ on $M = 10$ trials.

(Bottom) Results for contiguous LWCV.

Figure 7: Leave Future Out CV for MoCAP data on a single MoCAP sequence containing 1484 measurements. The scatter plots compare $-\ln p(x_{T'} | x_{[T]-o}; \Theta(\omega))$ (horizontal axis) with $-\ln p(x_{T'} | x_{[T]-o}; \hat{\Theta}_{IJ}(\omega))$ (vertical axis), with $o = \{T', T'+1, \ldots, T\}$, for some $T' \leq T$, for a five state HMM with Gaussian emissions (left), order 1 auto-regressive emissions (middle), neural auto-regressive emissions (right). The rightmost plot shows the number of parameters in each model. We vary $T'$ from 1337 to 1484 for Gaussian and AR(1) emissions. Since exact fits the NAR model are more expensive we only vary $T'$ between 1455 and 1484 for NAR(1). We find that ACV to be accurate. The NAR model which is an instance of a higher dimensional optimization problem, leads to approximations that are less accurate than the lower dimensional AR(0) and AR(1) cases.

Accurate LWCV for MoCAP Next, we present LWCV results on a 1,484 measurement long sequence extracted from the MoCAP dataset. We explore three variants of LWCV: i.i.d LWCV, contiguous LWCV, and a special case of contiguous LWCV: leave-future-out CV. Figures 6 and 7 present these results. We find that the IJ approximations again provide accurate approximations to exact CV. The performance deteriorates for contiguous LWCV when large chunks of the sequence are left out. Since large changes to the sequence result in large changes to the fit parameters, a Taylor series approximation about the original fit is less accurate. Also, for high dimensional models such as NAR(1) IJ approximations tend to be less accurate [Stephenson and Broderick, 2020], explaining the drop in LFOCV performance for the NAR(1) model.