Motivated by the analysis of torsion (dihedral) angles in the backbone of proteins, we investigate clustering of bivariate angular data on the torus $[-\pi, \pi) \times [-\pi, \pi)$. We show that naive adaptations of clustering methods, designed for vector-valued data, to the torus are not satisfactory, and propose a novel clustering approach based on the conformal prediction framework. We construct several prediction sets for toroidal data with guaranteed finite-sample validity, based on a kernel density estimate and bivariate von Mises mixture models. From a prediction set built from a Gaussian approximation of the bivariate von Mises mixture, we propose a data-driven choice for the number of clusters, and present algorithms for an automated cluster identification and cluster membership assignment. The proposed prediction sets and clustering approaches are applied to the torsion angles extracted from three strains of coronavirus spike glycoproteins (including SARS-CoV-2, contagious in humans). The analysis reveals a potential difference in the clusters of the SARS-CoV-2 torsion angles, compared to the clusters found in torsion angles from two different strains of coronavirus, contagious in animals.

1. Introduction. The structure of a protein is often summarized by the torsion (dihedral) angles formed along the backbone of the protein (Dill and MacCallum, 2012). A standard visual representation of the torsion angles is given by the Ramachandran plot, in which the sequence of $(\phi, \psi)$ torsion angles extracted from the backbone of a protein is plotted on $[-\pi, \pi) \times [-\pi, \pi)$ (Fig. 1). Due to the unique (circular) challenges in the relatively simple data structure, and to its importance in structural analysis of proteins and RNA, there have been a number of endeavors on density estimation, clustering and dimension reduction of data on the torus (Mardia, Taylor and Subramaniam, 2007; Mardia et al., 2008, 2012; Lennox et al., 2009; Eltzner, Huckemann and Mardia, 2018; Shapovalov, Vucetic and Dunbrack Jr, 2019; Gao et al., 2018; Nodehi et al., 2020). In particular, clusters in the torsion angles have been interpreted as local structures of the backbone of a protein, which determine the protein’s functions (Berg, Tymoczko and Stryer, 2002).

The circular nature of angles leads that the bivariate angle $(\phi, \psi)$ is on the torus, which may be embedded as an intrinsically two-dimensional manifold in $\mathbb{R}^3$, and can be cut-and-flattened as a square $T^2 = [-\pi, \pi) \times [-\pi, \pi)$ on $\mathbb{R}^2$ (cf. Fig. 1, Eltzner, Huckemann and Mardia, 2018). Therefore a point $(\pi - \epsilon, \pi)$ is closer to $(\pi - \epsilon, \pi - \epsilon)$ than $(\pi + 2\epsilon, -\pi + 2\epsilon)$ for some $\epsilon > 0$. Due to this geometric constraint, as we shall see in Section 3.1, most off-the-shelf clustering methods are not applicable, at least not without a proper adaptation.

In this article, we propose a novel approach for clustering on the torus, based on the conformal prediction framework (Vovk, Gammerman and Shafer, 2005; Lei, Robins and Wasserman, 2013; Lei et al., 2018). The conformal prediction framework is a method of constructing distribution-free prediction sets with finite-sample validity, but there has been no attempt of applications to circular variables in the literature. Following Lei, Robins and Wasserman (2013); Lei, Rinaldo and Wasserman (2015), we construct estimators of prediction sets for the protein torsion angles, based on a kernel density estimate on the torus (Di Marzio, Panzera...
and Taylor, 2011) and a finite mixture of bivariate von Mises distributions (Mardia, Taylor and Subramaniam, 2007; Mardia et al., 2012). The conformal prediction sets for toroidal data are described in Section 2. See Fig. 2 for the prediction sets obtained from the data displayed in Fig. 1. We construct several variants of prediction sets, based on which the proposed clustering of toroidal data is defined.

We propose to identify clusters by the connected components of prediction sets. We say $A_1, \ldots, A_K$ are connected components of (a closed set) $A \subset T^2$ if $A$ is the disjoint union of $A_k$’s and each $A_k$ cannot be divided into two disjoint non-empty closed sets. For a careful choice of conformity score (from which the prediction set $\hat{C}_n$ is estimated), the connected components are simply unions of ellipses, and an automated identification of clusters is possible. The resulting number $K$ of clusters depends on the choice of the level of the prediction set, and the hyperparameter used in fitting the mixture model. The problem of choosing $K$ is then transformed into the problem of setting the level and the hyperparameter, which are chosen by striking a balance between the desired coverage (the higher the better) and the volume of the prediction set (the smaller the better). We propose and compare two methods of cluster membership assignment. See Section 3 for the proposed approaches of clustering. Figure 2 demonstrates the results of the proposed clustering, in which three major clusters are identified.

The data shown in Figs 1 and 2 are torsion angles from Cryo-EM structures of SADS-CoV spike glycoproteins, previously analyzed in Yu et al. (2020). The swine acute diarrhea syndrome coronavirus (SADS-CoV) is a strain of coronavirus, and is known to be structurally similar to Rhinolophus bat coronavirus (HKU2) (Gong et al., 2017). We compare the clusters of SADS-CoV torsion angles with those of HKU2 and the severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2), a variant of RNA coronavirus (Walls et al., 2020), which is the cause of the COVID-19 pandemic (Gorbalenya et al., 2020; Chan et al., 2020). Section 4.1 is devoted to analyze the torsion angles from these coronavirus proteins, and we reveal a potential difference in the distributions of torsion angles of SARS-CoV-2 from those of SADS-CoV and HKU2.

The advantages of the proposed clustering framework are further highlighted in a simulation study (Section 4.2) by an empirical comparison with existing clustering methods on the torus. Our method takes the cyclic nature of toroidal data into account, and performs supe-
rior when the clusters are of irregular shape. Technical details and supplementary figures are supplied in the Appendix (the online supplementary material).

The proposed clustering approaches can be easily adapted for the usual multivariate data (in the Euclidean space) and can be extended to other types of manifold-valued data in, e.g., hyperspheres and higher-dimensional tori. In Section 5, we discuss related approaches in Euclidean spaces and point out future research directions.

1.1. Tools to handle data on the torus. Bivariate angular data can be understood as lying on the product of two unit circles and are naturally parameterized by angles in $T^2 = [−\pi, \pi) \times [−\pi, \pi)$ or in $[0, 2\pi) \times [0, 2\pi)$. We use the former, but our discussion is invariant to the choice of parameterization.

Let $x = (\phi_x, \psi_x), y = (\phi_y, \psi_y) \in T^2$. The angular subtraction is defined as $x \ominus y = \arg(e^{i(x-y)} := (\arg(e^{i(\phi_x-\phi_y)}), \arg(e^{i(\psi_x-\psi_y)})))$, in which operations are applied element-wise. (The range of the argument is $[−\pi, \pi]$.) As an example, $(\pi/2, 3\pi/4) \ominus (0, -3\pi/4) = (\pi/2, -\pi/2)$. A natural metric on the torus is $\rho(x, y) = \|x \ominus y\|_2 = \|y \ominus x\|_2 = [\{\arg(e^{i(\phi_x-\phi_y)})\}^2 + \{\arg(e^{i(\psi_x-\psi_y)})\}^2]^{1/2}$. For any $x, y \in T^2, \rho(x, y) \in [0, \sqrt{2}\pi]$. The toroidal distance function $\rho(x, y)$ can be used, for example, in defining the nearest neighbors of a point on $T^2$. Finally, given a set of points $X = \{X_1, \ldots, X_n\}$, the (sample) toroidal mean is defined by $m(X) = \arg\min_x \sum_{i=1}^n \rho^2(x, X_i)$, where the minimum is over $T^2$. The toroidal mean always exists, but may not be unique. Note that our parameterization of the torus with the metric $\rho$ makes the torus a flat torus (cf. O’neill, 2006).

2. Distribution-free prediction sets on the torus.
2.1. Conformal prediction framework. Suppose we observe an exchangeable sequence of random variables \( \{X_1, \ldots, X_n\} \) on the torus, where \( X_i = (\phi_i, \psi_i) \in \mathbb{T}^2 \). A level \( 1 - \alpha \) prediction set \( C_n = C_n(X_1, \ldots, X_n) \) satisfies, for new observation \( X_{n+1} \),
\[
P(X_{n+1} \in C_n) \geq 1 - \alpha,
\]
where the probability \( P \) is with respect to the exchangeable sequence \( \{X_1, \ldots, X_n, X_{n+1}\} \). Conformal prediction framework estimates \( C_n \) by introducing conformity scores. The conformity score \( \sigma_i = \sigma(X_{n+1}; X_i) \) measures the degree of conformity of \( X_i \) compared to the augmented sample \( X_{n+1} = \{X_1, \ldots, X_n, X_{n+1}\} \), and is defined for each \( X_i, i = 1, \ldots, n+1 \). For example, a conformity score defined by \( \sigma_i = \frac{1}{n+1} \sum_{j=1}^{n+1} \mathbf{1}_{\sigma_j \leq \sigma_{n+1}} \).

It can be shown that for any choice of conformity score \( \sigma \), the conformal prediction set \( C_n \) is a valid level \( 1 - \alpha \) prediction set, i.e. (1) holds, if \( X_{n+1} \) is exchangeable; see e.g. page 26 of [Vovk, Gammerman and Shafer (2005)] and Section 2 of [Lei, Robins and Wasserman (2013)]. A caveat here is that for a poorly-chosen conformity score \( \sigma \) the volume of \( C_n \) can be nontrivially larger than using more sensible choices of \( \sigma \). [Lei, Robins and Wasserman (2013)] has shown that if kernel density estimates are used for conformity scores, then \( C_n \) is efficient for data in \( \mathbb{R}^d \). In the next subsection, we adapt the idea of [Lei, Robins and Wasserman (2013)] for the prediction of toroidal data.

2.2. Prediction set by kernel density estimates on the torus. A natural population counterpart of prediction set \( C_n \) is the density level set \( L(t) = \{x \in \mathbb{T}^2 : p(x) \geq t\} \), where \( p(\cdot) \) is the density function of \( X_{n+1} \) on \( \mathbb{T}^2 \). For a given \( \alpha, t = t_\alpha \) is defined as the largest \( t \) satisfying \( P(X_{n+1} \in L(t)) \geq 1 - \alpha \). If the density function \( p \) is continuous on \( \mathbb{T}^2 \) and is not flat at \( \{x : p(x) = t_\alpha\} \), then it can be shown that the level set \( L(t_\alpha) \) has exact coverage and has the smallest volume among all level \( 1 - \alpha \) prediction sets [Polonik, 1997]. To be precise, let \( \mu(A) \) be the area of a subset \( A \subset \mathbb{T}^2 \), scaled by \( (2\pi)^2 \) so that \( \mu(\mathbb{T}^2) = 1 \). Then \( L(t_\alpha) = \arg\min_C \mu(C) \) where the minimum is over \( \{C \subset \mathbb{T}^2 : P(X_{n+1} \in C) \geq 1 - \alpha\} \).

We build a conformal prediction set \( C_n \) using a form similar to the density level set \( L(t_\alpha) \), in which \( p \) and \( t_\alpha \) are given by a kernel density estimate (kde) \( \hat{p}(\cdot) \) and conformity scores; see (5) below. This is done indirectly by using \( \hat{p}(\cdot) \) as the conformity score \( \sigma(X_{n+1}, \cdot) \).

We first discuss a kde on \( \mathbb{T}^2 \). Due to the doubly cyclic nature of the sample space \( \mathbb{T}^2 \), care is needed in defining a kde. For example, scalable kernels defined on a bounded domain such as the Epanechnikov kernel \( K(x) = \frac{3}{4}(1 - x^2)1_{|x| \leq 1} \) (or any generalization to \( \mathbb{T}^2 \)) does not integrate to 1 when scaled by \( x \mapsto x/s \), \( s > \pi \). Gaussian kernels suffer from the need of truncation on \( [-\pi, \pi] \) or of wrapping around \( \mathbb{T}^2 \). Instead, following [Di Marzio, Panzera and Taylor (2011)], we use a 2-product von Mises kernel (with a common concentration parameter \( \kappa \)): For \( x = (\phi, \psi) \in \mathbb{T}^2 \)
\[
K_\kappa(x) = \frac{e^{\kappa \cos(\phi)}}{2\pi I_0(\kappa)} \frac{e^{\kappa \cos(\psi)}}{2\pi I_0(\kappa)},
\]
where \( I_\nu \) is the modified Bessel function of the first kind of order \( \nu \). The von Mises density 
\[
(2\pi I_0(\kappa)^{-1} e^\nu \cos(\phi) \right)
plays the role of the normal distribution for circular statistics (Mardia and Jupp, 2000).

The kernel \((x)\) is a simple form of toroidal kernels. Kernel density estimates using toroidal kernels are known to enjoy some nice asymptotic properties (Di Marzio, Panzeri and Taylor, 2011). Using \((\)\), the kde of \( p(u) \) at \( u \in T^2 \) is \( \hat{p}_n(u) = \sum_{i=1}^{n} K_\kappa(u - X_i)/n \),
where the cyclic nature of \( T^2 \) is handled through the cosine function.

To define conformity scores, we use the kde based on the augmented data \( X_{n+1} \),
\[
\hat{p}_{n+1}(u) = \frac{1}{n+1} \sum_{i=1}^{n+1} K_\kappa(u - X_i)
\]
(4)
in which the unobserved \( X_{n+1} \) is replaced by \( X_{n+1} = x \), for an inspection point \( x \in T^2 \).

Setting \( \sigma_i = \hat{p}_{n+1}(X_i) \), the expression \( \xi_n(x) \) in the conformal prediction set \((\)\) becomes
\[
\xi_n(x) = \frac{1}{n+1} \left\{ 1 + \sum_{i=1}^{n} \frac{1}{\hat{p}_{n+1}(X_i)} \leq \hat{p}_{n+1}(x) \right\}.
\]
Observe that \( \xi_n(x) \in \{ \frac{1}{n+1}, \ldots, \frac{n}{n+1}, 1 \} \), which in turn leads that \( \pi_n(x) > \alpha \) is equivalent to \( \pi_n(x) > \tilde{\alpha} \), where
\[
\tilde{\alpha} = \frac{i_{n,\alpha}}{n+1}, \quad i_{n,\alpha} = \lfloor (n+1) \alpha \rfloor.
\]
To further simplify, let \( X_{(i)} \) (\( i = 1, \ldots, n \)) satisfy \( \hat{p}_{n+1}(X_{(i)}) \leq \hat{p}_{n+1}(X_{(i+1)}) \) for \( 1 \leq i \leq n-1 \). Then,
\[
C_n = \{ x \in T^2 : \xi_n(x) > \tilde{\alpha} \}
\]
(5)
\[
= \{ x \in T^2 : \hat{p}_{n+1}(x) \geq \hat{p}_{n+1}(X_{(in,\alpha)}) \}.
\]
The equation \((\)\) is verified in Appendix B.1. Note that \( C_n \) is not exactly a level set of a single function as the density estimate \( \hat{p}_{n+1} \) depends on \( x \). Moreover, to evaluate whether \( x \in C_n \) for many inspection points \( x \), the computation of \( \hat{p}_{n+1} \) and \( X_{(i)} \) is required for each \( x \). We provide two approximations of \((\)\) as level sets of \( \hat{p}_n \), which require a significantly less computation, in Appendix B.2.

As a demonstration, the prediction set \( C_n \) of level 0.9 is plotted in Fig. 2 (top left panel).

2.3. Prediction set by mixtures of bivariate von Mises. In this section, a mixture density and its variants are used as a conformity score \( \sigma \) in constructing prediction sets.

Our choice of the mixture model on the torus is composed of bivariate von Mises distributions (Mardia, Taylor and Subramaniam, 2007; Chakraborty and Wong, 2017). In particular, we use the sine variant of bivariate von Mises density (Singh, Hnizdo and Demchuk, 2002),
\[
f(x) = C \exp [\kappa_1 \cos(\phi - \mu_1) + \kappa_2 \cos(\psi - \mu_2) + \lambda \sin(\phi - \mu_1) \sin(\psi - \mu_2)],
\]
for \( x = (\phi, \psi) \in T^2 \), where \((\mu_1, \mu_2) \in T^2 \) is the location parameter, \( \kappa_1 > 0, \kappa_2 > 0 \) are concentration parameters and \( \lambda \), satisfying \( \lambda^2 < \kappa_1 \kappa_2 \), determines an association of two circular variables. The normalizing constant is given by
\[
\frac{1}{2m} \sum_{m=0}^{\infty} \left( \frac{\lambda^2}{4\kappa_1 \kappa_2} \right)^m I_m(\kappa_1) I_m(\kappa_2).
\]
If \( \lambda = 0 \), then \( f(\phi, \psi) \) is a product of two von Mises densities. For large concentrations (larger values of \( \kappa_1, \kappa_2 \), the density is well approximated by a bivariate normal density. For simplicity assume \( \mu_1 = \mu_2 = 0 \) for now. Then,
\[
f(x) \approx C' \exp \left( -\frac{1}{2} x^T \Sigma^{-1} x \right), \Sigma^{-1} = \left( \begin{array}{cc} \kappa_1 - \lambda & \lambda \\ \lambda & \kappa_2 \end{array} \right),
\]
(7)
where \( C' = |2\pi \Sigma|^{-1} = (\kappa_1 \kappa_2 - \lambda^2)/(2\pi)^2 \). The parameters \( \kappa_1, \kappa_2 \) and \( \lambda \) can be interpreted using (7): Writing \( \sigma_\phi^2, \sigma_\psi^2 \) and \( \rho \) for the variances of \( \phi \) and \( \psi \) and the correlation coefficient between them, respectively, we have \( \kappa_1^{-1} \approx \sigma_\phi^2(1 - \rho^2), \kappa_2^{-1} \approx \sigma_\psi^2(1 - \rho^2) \) and \( \lambda \approx \rho \sqrt{\sigma_\phi^2(1 - \rho^2)\sigma_\psi^2(1 - \rho^2)} \). We require \( \lambda \in (-\sqrt{\kappa_1 \kappa_2}, \sqrt{\kappa_1 \kappa_2}) \), under which condition the bivariate von Mises density is unimodal (Theorem 3, Mardia, Taylor and Subramaniam, 2007).

For a \( J = 1, 2, \ldots, \) a \( J \)-mixture density for the toroidal data is given by \( p(x) = \sum_{j=1}^{J} \pi_j f_j(x) \), where \( \pi_j \)'s are mixing probabilities (\( \sum_j \pi_j = 1 \)) and \( f_j(\cdot) \) is the bivariate von Mise density with parameters \( \theta_j := (\mu_{1j}, \mu_{2j}, \kappa_{1j}, \kappa_{2j}, \lambda_j) \).

Maximum likelihood estimates of \( \pi_j, \theta_j \) and \( f_j(\cdot) \) based on \( \mathcal{X}_n = \{X_1, \ldots, X_n\} \) are denoted by \( \hat{\pi}_j, \hat{\theta}_j \) and \( \hat{f}_j(\cdot) \), respectively, which provide a natural definition of \( \hat{p}_n^M(\cdot) \) (M stands for mixture models). These estimates are numerically obtained by an EM algorithm. The details of the estimation procedure are discussed in Appendix A.1. Let \( \hat{p}_{n+1}^M \) be defined similarly for an augmented data set \( \mathcal{X}_{n+1} \). Then

\[
C_n^M = \{ x \in \mathbb{T}^2 : \hat{p}_{n+1}^M (X_i) \geq \hat{p}_{n+1}^M (X_{i(n_n+1)}) \}
\]

is a valid level \( 1 - \alpha \) prediction set constructed by setting \( \sigma_i = \hat{p}_{n+1}^M (X_i) \); compare with equation (5) and the preceding discussion.

Computing \( \hat{p}_{n+1}^M \) for many \( x \) values of the augmented data \( \mathcal{X}_{n+1} \) with \( X_{n+1} = x \) is not practical. Instead, we use a sample splitting strategy, called \textit{inductive conformal prediction} in the literature of conformal prediction (Vovk, Gammerman and Shafer, 2005; Lei, Rinaldo and Wasserman, 2015). The idea is to randomly split \( \mathcal{X}_n \) into \( \mathcal{X}_1 \) of size \( n_1 \) and \( \mathcal{X}_2 \) of size \( n_2 \) (\( n_1 + n_2 = n \)). The first part \( \mathcal{X}_1 \) is used in the estimation of \( \hat{p} \), resulting in \( \hat{p}_{n_1}^M \) (denoted by \( \tilde{p} \) hereafter for notational simplicity). The conformity scores are computed for the second part \( \mathcal{X}_2 = \{X_{n_1+1}, \ldots, X_n\} \): \( \sigma_i = \hat{p}(X_{n_1+i}) \) (\( i = 1, \ldots, n_2 \)), then sorted to \( \sigma(1) \leq \ldots \leq \sigma(n_2) \). A level \( 1 - \alpha \) inductive conformal prediction set is the level set of \( \tilde{p} \):

\[
\tilde{C}_n = \{ x \in \mathbb{T}^2 : \tilde{p}(x) \geq \sigma_{(i_{n_2,\alpha})} \}, \quad i_{n_2,\alpha} = \lfloor (n_2 + 1) \alpha \rfloor.
\]

As the computation of \( \tilde{p} \) is required only once in (8), evaluating the inductive prediction set has a huge computational advantage over evaluating \( C_n^M \). It is shown in Lei, Rinaldo and Wasserman (2015) that inductive conformal prediction set also enjoys the distribution-free finite-sample validity, i.e. \( P(\mathcal{X}_{n+1} \in \tilde{C}_n) \geq 1 - \alpha \). We emphasize that \( \tilde{C}_n \) is valid not only when the mixture density estimate \( \hat{p}_{n_1}^M \) is used in \( \sigma_i \) but also for \textit{any} choices of conformity scores. Thus the variants of \( \tilde{C}_n \) we discuss below also have the finite-sample validity.

Suppose that the components in \( \tilde{p} \) are well separated, then

\[
\tilde{p}(x) = \sum_{j=1}^{J} \hat{\pi}_j \hat{f}_j(x) \approx \hat{p}^{\text{max}}(x) = \max_j \hat{\pi}_j \hat{f}_j(x).
\]

This suggests setting the conformity scores to \( \sigma_i = \hat{p}^{\text{max}}(X_{n_1+i}) \), which in turn leads to a valid inductive conformal prediction set (even if the components are not separated).

Another variant is obtained when considering the normal approximation to the bivariate von Mises (7). The log-max-density \( \log(\hat{p}^{\text{max}}(x)) = \max_j \{ \log(\hat{\pi}_j) + \log(\hat{f}_j(x)) \} \) is approximated by \( \hat{e}(x)/2 + \text{constant} \), where

\[
\hat{e}(x) = \max_j \hat{e}_j(x),
\]

\[
\hat{e}_j(x) = -(x \odot \hat{\mu}(j))^T \hat{\Sigma}_j^{-1} (x \odot \hat{\mu}(j)) + 2 \log \left( (\hat{\kappa}_1 \hat{\kappa}_2 - \hat{\lambda}_j^2) \hat{\pi}_j \right).
\]

Here, \( \hat{\mu}(j) = (\hat{\mu}_{1j}, \hat{\mu}_{2j}) \), and \( \hat{\Sigma}_j^{-1} \) is given in (7) with the parameters replaced by the \( j \)th component estimates. The “\( x \odot y \)” notation refers to the angular subtraction, defined in Section 1.1. Any level set of \( \hat{e}(\cdot) \) is a union of ellipses on \( \mathbb{T}^2 \), as shown in Lemma 2.1.
Lemma 2.1. Let \( \hat{e}(x) \) be defined in (9), and let
\[
c_j = 2 \log \left( (\hat{\kappa}_{1j}\hat{\kappa}_{2j} - \hat{\lambda}^2_j)^{\hat{\pi}_j} \right)
\]
and
\[
\hat{E}_j(t) = \{ x \in \mathbb{T}^2 : (x \ominus \hat{\mu}_{(j)})^T \hat{\Sigma}^{-1}_j (x \ominus \mu_{(j)}) \leq c_j - t \}.
\]
Then for any \( t \in \mathbb{R} \), \( L^\hat{e}(t) := \{ x \in \mathbb{T}^2 : \hat{e}(x) \geq t \} = \bigcup_{j=1}^t \hat{E}_j(t) \).

Proof of Lemma 2.1. Write \( \hat{e}(x) = \max_j \hat{e}_j(x) \), and observe that \( \hat{e}_j(x) \geq t \) is equivalent to the condition for \( x \in \hat{E}_j(t) \). Then, \( \{ x \in \mathbb{T}^2 : \max_j \hat{e}_j(x) \geq t \} = \{ x \in \mathbb{T}^2 : \hat{e}(x) \geq t \} \) for some \( j \). \( \hat{E}_j(t) \) degenerates to an empty set.

Each \( \hat{E}_j(t) \) is an ellipse*, provided that \( t < c_j \). Otherwise, \( \hat{E}_j(t) = \emptyset \) for \( c_j - t \leq 0 \). Note that as \( t \) increases, the ellipses \( \hat{E}_j(t) \) of \( L^\hat{e}(t) \) become simultaneously smaller, and the number of ellipses involved decreases (as some \( \hat{E}_j(t) \) degenerates to an empty set).

Setting \( \sigma_j = \hat{e}(X_{n_1+i}) \) (which are then ordered to satisfy \( \sigma(1) \leq \cdots \leq \sigma(n_x) \)), the inductive conformal prediction set is
\[
\hat{C}_n^\sigma = \{ x \in \mathbb{T}^2 : \hat{e}(x) \geq \sigma_{(n_2,n)} \} = L^\hat{e}(\sigma_{(n_2,n)}).
\]

An advantage of using \( \hat{C}_n^\sigma \) is its simple form (the union of ellipses), which is computationally handy when we consider clustering in Section 3.

We denote the inductive prediction sets by \( \hat{C}_n^{\text{max}} \) if \( \sigma(\cdot) = \hat{p}(\cdot) \) (8), \( \hat{C}_n^{\text{max}} \) if \( \sigma(\cdot) = \hat{p}^{\text{max}}(\cdot) \), and \( \hat{C}_n^s \) if \( \sigma(\cdot) = \hat{e}(\cdot) \) (10). The prediction set \( \hat{C}_n^s \) is plotted in Fig. 2 for the SADS-CoV torsion angles, further discussed in Section 4.1. (All prediction sets are displayed in the Appendix Fig. C.2.) An inductive prediction set \( \hat{C}_n^\text{kde} \) given by the kde is defined similarly.

3. Clustering on the torus.

3.1. Clustering on the torus: an overview. Due to the cyclic nature of data on \( \mathbb{T}^2 \), recklessly applying off-the-shelf clustering methods can result in unstable and low-quality clustering results. For example, the result of a naive k-means clustering applied to \( x_i \in \mathbb{T}^2 \) is different from the clustering results from the data \( x'_i := x_i + (\pi, \pi) \in [0, 2\pi)^2 \). However, since the data set \( \{ x'_i \} \) is simply translated from \( \{ x_i \} \), the clustering result should be identical. Moreover, an apparent cluster near the border of the square \( [-\pi, \pi] \times [-\pi, \pi] \) is split into two or more clusters, as exemplified in the top left panel of Fig. 3. Some existing work on clustering toroidal data (Kountouris and Hirst, 2009) suffers from the same problem.

We briefly review adaptations of a few popular clustering methods for use on the torus.

The k-means clustering is by far the most popular off-the-shelf clustering algorithm, obtained by recursively partitioning observations and computing cluster centers. A straightforward adaptation of the k-means for data on the torus is given by using the toroidal distance \( \rho(x,c) \) (in partitioning) and the toroidal mean \( m(X) \) (for cluster centers). An approximation of this intrinsic k-means algorithm was considered in Gao et al. (2018). In contrast, an extrinsic k-means algorithm uses the ambient space for \( \mathbb{T}^2 \), in which each \( \mathbb{T} = [-\pi, \pi] \) is embedded as the unit circle in \( \mathbb{R}^2 \). Each \( x_i = (\phi_i, \psi_i) \in \mathbb{T}^2 \) is transformed

*Precisely, the set \( \{ x \in \mathbb{T}^2 : (x \ominus \mu)^T S^{-1} (x \ominus \mu) \leq 1 \} \) is an ellipse for small enough \( S \); for large \( S \), the set is the intersection of an ellipse with the square of width \( 2\pi \) centered at \( \mu \). In our analysis, \( S = \max \{ c_j - t, 0 \} \hat{\Sigma}_j \) is typically small.
to $x'_i = (\cos \phi_i, \sin \phi_i, \cos \psi_i, \sin \psi_i) \in \mathbb{R}^4$, then a usual k-means algorithm is applied to $x'_i$s.

This extrinsic k-means algorithm is computationally advantageous as it uses the vector operations in $\mathbb{R}^4$, and any variant of the k-means algorithm can be used as well. In Fig. 3, we demonstrate the result of the extrinsic k-means clustering for the protein structure data, with initial values given by the k-means++ algorithm (Arthur and Vassilvitskii, 2007).

A number of clustering algorithms depend only on the pairwise distances (or similarity measures) of data (Xu and Tian, 2015). The partitioning around medoids (PAM) algorithm of Kaufman and Rousseeuw (2009) [originally proposed in 1990], its variants (Van der Laan, Pollard and Bryan, 2003), and hierarchical clustering methods (Murtagh and Contreras, 2012, 2017) are prominent examples. All of these clustering methods are readily available for data on the torus with pairwise toroidal distances \( \{\rho(x_i, x_j) : 1 \leq i < j \leq n\} \) as an input.

Gaussian mixture models are another popular and well-developed class of clustering methods for vector-valued data; see, e.g., Scrucca et al. (2016). A toroidal adaptation of mixture models is given in Section 2.3 in which we used bivariate von Mises mixture models (Mardia, Taylor and Subramaniam, 2007). Using a fitted mixture model $\hat{p}(\cdot) = \sum_j \hat{\pi}_j \hat{f}_j(\cdot)$, a soft clustering of an inspection point $x$ is the probability of the cluster membership $Y$, $\hat{p}(Y = j \mid X = x) = \hat{\pi}_j \hat{f}_j(x) / \hat{p}(x)$ for $j = 1, \ldots, K$. A hard clustering of $x$ is simply $\arg \max_j \hat{\pi}_j \hat{f}_j(x)$.

Examples in Fig. 3 suggest that when the apparent clusters are of irregular shapes (rather than having elliptical shapes), all of the methods above provide unsatisfactory clustering results. For such cases, a density level set clustering might be useful (Hartigan, 1975; Cheng, 1995). For a multi-modal density $p$, the level set $L(t) = \{x \in T^2 : p(x) \geq t\}$ for appropriately chosen $t$ separates each mode from the others. However, it is unclear what should be an actual clustering function that assigns a cluster label for each $x \in T^2$. A naive approach of partitioning around medoids works poorly when the clusters (or the connected components of $L(t)$) are of irregular shapes.
In the next subsection, we develop a conformity-score level set clustering for the torus, and show that a sensible and automated cluster-label assignment can be done by employing a carefully chosen conformity score.

3.2. Clustering by conformal prediction sets. A conformal prediction set $C_n$ on the torus developed in Section 2 provides a natural clustering of $\mathbb{T}^2$, where a cluster is given by a connected component of $C_n$. If an inductive conformal prediction set (8) or (10) is used instead of $C_n$, then the corresponding clustering is equivalent to a density level set clustering (Hartigan, 1975), except that the density function $p(x)$ is replaced by the conformity score $\sigma(x)$, which is not necessarily a density estimate.

We point out that a prediction set (denoted by $\hat{C}$, representing either $C_n$ (5), $C_{n,\text{mix}}$ (8), $C_{n,\text{mixture}}$ (10), or $C_{n,\text{dese}}$) depends on the choice of level $1 - \alpha$ and a hyperparameter (concentration $\kappa$ for a kde-based prediction set, or the number $J$ of mixture components for a mixture-model-based prediction set). The inherent problem of choosing the number $K$ of clusters is substituted by the problem of selecting $\alpha$ and the hyperparameter. Our approach for the selection of level and hyperparameter will be discussed in Section 3.3. For now, let us assume that a prediction set is given for a prespecified choice of $\alpha$ and $\kappa$ (or $J$).

Identification of connected components. For any given prediction set $\hat{C}$, its connected components and the number of distinct components $K$ can be computed algorithmically using a fine grid on $\mathbb{T}^2$. Our visual illustrations of $\hat{C}$ in Fig. 2 (top row) are indeed given by evaluating $\mathbf{1}_{x \in \hat{C}}$ for $x \in \mathbb{T}^2$, where $T = \{\pi(2t - 1) : t = 1/100, 2/100, \ldots, 1\}$. To identify all grid points of a connected component containing $x$, one may use a flood fill algorithm, i.e., recursively identifying neighborhoods of $x$ in $\hat{C}$, which is then applied to all unidentified points in $\hat{C}$ to assign $x$ with labels distinct for each connected component. A drawback of such an algorithmic approach is that its accuracy deteriorates when the grid $T$ is coarse.

When the elliptical prediction set $\hat{C}_{n,e}^a$ is used, an exact identification of the connected components is possible. Recall that $\hat{C}_{n,e}^a = \bigcup_{j=1}^J \hat{E}_j$, where $\hat{E}_j = \hat{E}_j(\sigma_{(i,e,a)})$, as defined in Lemma 2.1. In this case, the connected components are exactly unions of ellipses. To identify the connected components, we create an adjacent matrix $A$ (of size $J \times J$) whose $(i,j)$th element is 1 if $\hat{E}_i \cap \hat{E}_j \neq \emptyset$, 0 otherwise. (We discuss complications in testing whether two toroidal ellipses intersect in Appendix A.2.) The adjacent matrix $A$ gives rise to an undirected graph (where nodes are labeled $1, \ldots, J$) whose connected components are easily found by a simple breadth first search. If $\hat{E}_j = \emptyset$, then the corresponding node $j$ is removed from the graph. Denoting the connected components by the graph by the node indices $\ell_1, \ldots, \ell_K \subset \{1, \ldots, J\}$, the (connected components) of $\hat{C}_{n,e}^a$ are

$$\mathcal{E}_k = \bigcup_{j \in \ell_k} \hat{E}_j, \quad k = 1, \ldots, K.$$ 

Note that $\mathcal{E}_k \cap \mathcal{E}_{k'} = \emptyset$ for $k \neq k'$, $\hat{C}_{n,e}^a = \bigcup_{k=1}^K \mathcal{E}_k$, and the number of clusters is $K$.

As an illustration, Fig. 4 displays a toy data set on $\mathbb{T}^2$, overlaid with $\hat{E}_j$’s, the union of which provides a 90% prediction set. For this data set, $\mathcal{E}_1 = \hat{E}_1$, $\mathcal{E}_2 = \hat{E}_2 \cup \hat{E}_3$ and there are $K = 2$ clusters.

Cluster assignment. Every $x \in \hat{C}$ has a natural cluster membership assignment, given by the label of a connected component it belongs to.

For $x \notin \hat{C}$, a natural approach of membership assignment is to find the “closest” cluster. When the closeness is defined by the toroidal distance between $x$ and a connected component $\mathcal{E}_k$, finding the closest cluster can be handled by a grassfire transformation (Blum, 1967). However, this approach has a statistical drawback. To motivate, consider two components of a bivariate von Mises (vM2) mixture: One is approximately given by a vM2 with parameters $\theta_1 = (0, 0, 50, 3, 0)$, the other by $\theta_2 = (-\pi, 0, 3, 50, 0)$. Ellipses corresponding to these
two vM2 components are shown in Fig. 4. There, \( \hat{C}_{en} \) is obtained by a 4-mixture of bivariate von Mises with level \( 1 - \alpha = 90\% \). Here, \( \hat{E}_4 = \emptyset \) is not shown. There are \( K = 2 \) connected components, and the point \( x \) is assigned to the second cluster \( \hat{E}_2 \) by \( A_e(x) = 2 \) (11).

Suppose that the clustering is based on the mixture model fitting and that \( X \) follows the \( J \)-mixture of bivariate von Mises distributions and let \( Y \in \{1, \ldots, J\} \) be the unobservable component membership of \( X \). Given connected component indices \( \ell_1, \ldots, \ell_K \subset \{1, \ldots, J\} \), an \( x \in \mathbb{T}^2 \) is assigned to cluster \( \hat{y}(x) = \arg\max_k \hat{P}(Y \in \ell_k | X = x) \), where \( \hat{P}(Y \in \ell_k | X = x) \) is proportional to \( \sum_{j \in \ell_k} \hat{\pi}_j \hat{f}_j(x) \). Denote this assignment rule as \( A_p \) so that for any \( x \in \mathbb{T}^2 \),

\[
A_p(x) = \arg \max_k \sum_{j \in \ell_k} \hat{\pi}_j \hat{f}_j(x).
\]

If components are well-separated, then \( \sum_{j \in \ell_k} \hat{\pi}_j \hat{f}_j(x) \approx \max_{j \in \ell_k} \hat{\pi}_j \hat{f}_j(x) \) and the maximum over \( k \) is well approximated by the maximum over \( k \) of \( \max_{j \in \ell_k} \hat{e}_j(x) \); see (9). Thus, an alternative cluster assignment rule, especially for use with \( \hat{C}^n_{en} \), is given by assigning the cluster label of \( x \) to \( k \) if \( \arg \max_j \hat{e}_j(x) \in \ell_k \):

\[
A_e(x) = k \text{ if } \arg \max_j \hat{e}_j(x) \in \ell_k, \ k = 1, \ldots, K.
\]

Note that for the mixture-based \( \hat{C}^n_{mix} \) (8), the assignment rule \( A_p \) guarantees that for \( x \in \hat{C}^n_{mix} \), \( A_p(x) \) is the label of a connected component it belongs to. Likewise, for the ellipse-based \( \hat{C}^n_{en} \), \( A_e(x) \) has the correct label.

Another approach is simply creating a new label, representing outliers, for all \( x \notin \hat{C} \):

\[
A_o(x) = \begin{cases} 
k, & x \in \mathcal{E}_k, \ k = 1, \ldots, K; \\
\text{"outliers"}, & \text{otherwise}.
\end{cases}
\]
3.3. **Selection of level, hyperparameters, and the number of clusters.** Our proposed clustering, introduced in Section 3.2, depends on the choice of level $1 - \alpha$ and a hyperparameter (the number of mixture components $J$ if using $\hat{C}_n^e$ or the concentration parameter $\kappa$ if using $\hat{C}_n^{kde}$). Without losing generality, we focus on the clustering based on $\hat{C}_n^e$ and present a data-driven approach of choosing $\alpha$ and $J$. Write $\hat{C}_n^e = \hat{C}_n^e(\alpha, J)$ as it depends on $\alpha$ and $J$. The number of clusters $K$ is just the number of connected components of $\hat{C}_n^e(\alpha, J)$.

As a prediction set, a desired level $1 - \alpha$ of $\hat{C}_n^e$ may be predefined. For a fixed $\alpha$, which choice of $J$ provides the best prediction set? This question was answered for a kde-based prediction set in Lei, Robins and Wasserman (2013). Since the coverage probability of $\hat{C}_n^e(\alpha, J)$ is guaranteed to exceed $1 - \alpha$, a prediction set with the smallest volume is desirable. The optimal choice of $J$, given an $\alpha$, is

$$J_\alpha = \arg\min_J \mu\left(\hat{C}_n^e(\alpha, J)\right),$$

where the area $\mu(A)$ of $A \subset \mathbb{T}^2$ is scaled to satisfy $\mu(\mathbb{T}^2) = 1$.

With clustering in mind, there is no predetermined level $1 - \alpha$, but the number $J$ of mixture components, or equivalently the fitted $J$-mixture bivariate von Mises model $(\hat{\pi}_j, \hat{f}_j(\cdot))$, may be given. The choice of $\alpha$ affects the number of clusters $K$ more gravely than the choice of $J$. As the coverage $1 - \alpha$ becomes larger, more components are connected to each other, which results in a smaller number of clusters. Ideally, one would choose as large coverage $1 - \alpha$ as possible and, simultaneously, wish for as small volume of the prediction set $\hat{C}_n^e(\alpha, J)$ as possible. However, there is a trade-off between large coverage (or, equivalently, small $\alpha$) and small $\mu(\hat{C}_n^e(\alpha, J))$. We propose to choose $\alpha$ by

$$\alpha_J = \arg\min_\alpha \alpha + \mu\left(\hat{C}_n^e(\alpha, J)\right).$$

Choosing $\alpha$ by $\alpha_J$ is equivalent to finding an “elbow” of the graph of the function $\alpha \mapsto \mu(\hat{C}_n^e(\alpha, J))$. As an attempt of interpretation, we note that increased coverage (by choosing $\alpha < \alpha_J$) leads for the volume of prediction sets to increase sharply; on the other hand, by reducing the volume ($\alpha > \alpha_J$), the coverage probability $1 - \alpha$ decreases fast. Setting the coverage at $1 - \alpha_J$ strikes a balance.

To choose both $\alpha$ and $J$ altogether, (13) and (14) are combined to

$$\left(\alpha_J, J\right) = \arg\min_{\alpha, J} \alpha + \mu\left(\hat{C}_n^e(\alpha, J)\right),$$

which is equivalent to finding the most lower-left point of $\{(\alpha, \mu(\hat{C}_n^e(\alpha, J))) : \alpha \in (0, 1), J = 1, \ldots, J_{\max}\}$.

### 4. Clustering protein torsion angles.

4.1. **Analysis of torsion angles from coronavirus protein structures.** We analyze torsion angles from Cryo-EM structures of SADS-CoV, HKU2, and SARS-CoV-2 (coronavirus) spike glycoproteins (Yu et al., 2020; Walls et al., 2020). Coronaviruses are a large group of viral pathogens, and pose severe threats to world healths when transmitted to humans. Severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2) is a single-stranded RNA coronavirus, and is contagious in humans (Gorbalenya et al., 2020; Chan et al., 2020). The swine acute diarrhea syndrome coronavirus (SADS-CoV), widespread in 2017 among commercial pigs in China, is found to share identical sequences up to 95% with Rhinolophus bat coronavirus (HKU2) (Gong et al., 2017). In this analysis, we confirm that SADS-CoV and HKU2 have almost identical protein backbone torsion angles, but SARS-CoV-2 has different sets of torsion angle clusters when compared to SADS-CoV.
Conformal prediction sets $\hat{C}_{kde}^n(\alpha, \kappa)$, the boundaries of which are displayed as gray curves, for the SADS-CoV torsion angles.

We use the $(\phi, \psi)$ torsion angles extracted from SADS-CoV to illustrate our approaches of prediction sets and clustering, and use the respective torsion angle data from HKU2 and SARS-CoV-2 as a validation set. The clustering of torsion angles from SARS-CoV-2 turns out to be different from the clustering for SADS-CoV. The data are available to public in the worldwide Protein Data Bank (PDB) with codes 6VXX, 6M15 and 6M16. The R packages bio3d (Grant et al., 2006) was used to extract the data from the PDB.

The SADS-CoV protein structure is available from sequence 20 to 998. The torsion angle $\phi$ is not defined for the first location, and $\psi$ for the last location. Excluding the first and last locations and other missing values, the SADS-CoV data set we analyze is the sequence of $(\phi_i, \psi_i) \in \mathbb{T}^2$ of length $n = 964$. Similar preprocessing gives HKU2 and SARS-CoV-2 torsion angles of sizes 964 and 824. Note that each of these three torsion angles data sets is sequentially observed along the backbone of a protein. The torsion angles are thus in fact serially correlated, and the assumption of exchangeability may not hold in general. Nevertheless, inspecting the scatterplot of the $(\phi_i, \psi_i)$ angles has been useful in understanding a protein structure (Mardia, Taylor and Subramaniam, 2007; Eltzner, Huckemann and Mardia, 2018). Our analysis only uses the scatter of torsion angles, which can then be assumed independent.

Prediction sets and the choice of tuning parameters. Since the sample size $n = 964$ is not small, it makes sense to construct the inductive conformal prediction sets. We demonstrate the use of conformity scores based on kernel density estimates and the estimated bivariate von Mises mixtures. The concentration parameter $\kappa > 0$ plays the role of the reciprocal of the bandwidth in the usual kernel density estimation. For large $\kappa$, the prediction set $\hat{C}_{kde}^n(\alpha, \kappa)$ is prone to catch random fluctuations in the data set, as exemplified for $\hat{C}_{kde}^n(0.1, 100)$, shown in the bottom right panel of Fig. 5. The difference of $\hat{C}_{kde}^n(\alpha, \kappa)$ according to varying $\kappa$ is less severe if the coverage $1 - \alpha$ is high. As an instance, the 98% prediction sets shown in
the left panels of Fig. 5 are similar to each other, in spite of different choices of \( \kappa = 25, 100 \). Choosing \((\alpha, \kappa) = (0.1, 25)\) gives a nice separation of three connected components of the prediction sets (top right panel).

A bivariate von Mises mixture distribution with \( J = 10 \) components is also fitted to the data, with the restriction of \( \lambda = 0 \). (Other choices of \( J \) are considered later.) Prediction sets are obtained from this mixture model at various nominal level \( 1 - \alpha \). For cluster identification and assignment, the prediction set \( \hat{C}_n^e \), consisting of the union of ellipses, is convenient in identifying the connected components. To inspect the effect of varying \( \alpha \), we have plotted \( \hat{C}_n^e = \bigcup_{j=1}^J \hat{E}_j(\sigma_{(i+\alpha)}^{(j)}) \) and each of the ellipses (\( \hat{E}_j \)'s) for a few choices of \( \alpha \) in Fig. 6. When the coverage \( 1 - \alpha \) is large, the ellipses involved are large as well, resulting in a smaller number of connected components. As \( \alpha \) increases, the radii of all ellipses decrease, which results in a connected component either to be divided into two components or to disappear. For example, the largest connected component (labeled 1) for the \( \alpha = 0.02 \) case is divided into two connected components (labeled 1 and 2) for the \( \alpha = 0.05 \) case; see the top two panels of Fig. 6. On the other hand, comparing \( \alpha = 0.05 \) with 0.1 (bottom left panel), the connected component labeled 4 at \( \alpha = 0.05 \) disappears at \( \alpha = 0.1 \), as the ellipse becomes an empty set.

For each choice of \( \alpha \), the number \( K \) of clusters are counted as the number of connected components and the cluster membership \( A_e(x) \) (11) is computed for all torsion angles \( x \) in the SADS-CoV protein structure, also shown in Fig. 6. It is evident that the clustering results depend on the choice of \( \alpha \), and the hyperparameter \( J \) used in fitting the mixture model.

If there is a desired level of coverage \( 1 - \alpha \), then the best hyperparameter \( J \) (or \( \kappa \)) can be chosen to minimize the volume \( \mu(C) \) of the corresponding prediction set \( C \) (13). For the SADS-CoV torsion angles, the graph of \((\kappa, \mu(C))\) is roughly convex (as shown the top left panel of Fig. 7), leading that a choice of \( \kappa \) near 30 is a stable choice. Note that the kde-based prediction set \( \hat{C}_{n\text{kde}}^e(\alpha, \kappa) \) is continuous with respect to both \( \alpha \) and \( \kappa \). On the other hand,
the volume of the mixture-model based prediction set $\mu\{\hat{C}_n(\alpha, J)\}$ has a large fluctuation over increasing $J$ (see the top right panel of Fig. 7), indicating that estimating the mixture model with a large number of components from a sample of size $n_1 = 482$ might lead to unstable estimates. For the SARS-CoV-2 torsion angles, the graph of $(\alpha, \mu\{\hat{C}_n(\alpha, J)\})$ is roughly convex and smooth (see Appendix Fig. C.4). Nevertheless, in view of the criterion $\nu(\alpha, J) := \alpha + \mu(\hat{C}_n)$ (15), changes due to the level is more substantial than the changes over $J$. This is illustrated in the bottom panel of Fig. 7, where the values of $\nu(\alpha, J)$ are evaluated and plotted for various combinations of $\alpha \in (0, 0.5)$ and a few choices of $J \in \{3, \ldots, 35\}$. The criterion was minimized to be $\nu(\hat{\alpha}, \hat{J}) \approx 0.241$ at $\hat{\alpha} \approx 0.079$, $\hat{J} = 12$, at which more than 92% of the sample is bound to lie in the prediction set $\hat{C}_n$ whose volume is $100\mu(\hat{C}_n)\% \approx 14.1\%$ of the torus. Here, $\hat{C}_n = \hat{C}_n$. The level $\hat{\alpha}$ and the number of components $\hat{J} = 12$ are used in the subsequent analysis of clustering for SADS-CoV torsion angles. A similar analysis done for SARS-CoV-2 leads $\hat{\alpha} \approx 0.10$, $\hat{J} = 22$.

**Clustering and a post analysis.** For each set of SADS-CoV and SARS-CoV-2 torsion angles, we obtained $K = 3$ clusters $E_k$ for the SADS-CoV ($K = 6$ clusters for the SARS-CoV-2) and evaluated the cluster membership $A_e(x)$ (11) for each data set. See Fig. 8.

The protein structure is known to follow strict geometric rules. The three larger clusters (shown in both cases) correspond to the well-known shapes of protein structures (Walther and Cohen, 1999; Lovell et al., 2003). In particular, The cluster labeled 1 in the top panel of Fig. 8 is related to the right-handed $\alpha$-helix, Cluster 2 is to the $\beta$-sheet and Cluster 3 is to
Fig 8. Clustering results for SADS-CoV torsion angles (top) and SARS-CoV-2 torsion angles (bottom).

the left-handed $\alpha$-helix. In the bottom panel of Fig. 8 for the SARS-CoV-2, three additional smaller clusters are shown. It appears that Cluster 3 (filled squares) corresponds to a less common conformation called $\gamma$-turns (cf. Fig. 1 of Lovell et al., 2003). Clusters 5 and 6 of the SARS-CoV-2 do not overlap with any torsion angles of SADS-CoV, indicating a potential difference in protein structures of the two coronaviruses.

Empirical coverage of prediction regions. The SADS-CoV protein structure is known to be nearly identical to the HKU2 (Yu et al., 2020). On the other hand, SARS-CoV-2 is a different strain of coronavirus, whose protein backbone structure differs from the others. The prediction sets $\hat{C}_n$, based on which the proposed clustering is conducted, are theoretically valid, i.e. $P(x \in \hat{C}_n) \geq 1 - \alpha$, for all $\alpha \geq 0$, for any $n$ and for any choice of prediction set $\hat{C}_{\text{kde}}^n$, $\hat{C}_{\text{e}}^n$, $\hat{C}_{\text{mix}}^n$ and $\hat{C}_{\text{max}}^n$. We empirically confirm that the coverage of $\hat{C}_n$ (estimated from the SADS-CoV torsion angles) meets the nominal level $1 - \alpha$ for the SADS-CoV and HKU2 torsion angles. In the top two panels of Fig. 9, the empirical coverage probability $P(x \in \hat{C}_n)$
is near the nominal level $1 - \alpha$ and is generally above the lower bound of a 98% pointwise confidence interval for $P(x \in \hat{C}_n \mid \hat{C}_n)$, for most values of $\alpha \in (0, 0.2)$.

For the SARS-CoV-2 angles, shown in the bottom panel of Fig. 9, the coverage of the SADS-CoV-based prediction sets is below the 98% pointwise confidence lower bound for higher levels $1 - \alpha > 0.9$. This indicates again a potentially different backbone structure of the SARS-CoV-2 protein. Caution is needed in making a conclusion since the confidence limits are obtained for the given prediction set. In Appendix B.4, we show that the variance due to the estimation of $\hat{C}_n$ is larger than the variance due to the evaluation of the coverage, from a simulated data set of Section 4.2.

The hyperparameters for the prediction sets in Fig. 9 are chosen to be $\hat{\kappa} = 30$ and $\hat{J} = 12$ based on the criterion (15). Other choices of $\kappa$ and $J$ lead similar results.

4.2. Clustering artificial toroidal data. The proposed clustering procedure is tested upon two artificial data on the torus. We empirically compare the clustering performances of our proposal with existing methods in Section 3.1.

The two artificial data are each sampled from the following models:

- Model I: The dataset of size $n = 270$ is sampled from a mixture of $K = 5$ clusters, where three clusters are sampled from bivariate normal distributions (with sizes 70, 50, 50) and the other two are each sampled from the uniform distribution on a rectangle defined on $\mathbb{R}^2$ (each with size 50), then wrapped onto the torus.
- Model II: The dataset of size $n = 500$ is sampled from a mixture of $K = 3$ clusters, where the first cluster is sampled from a spherical normal distribution with size $n_1 = 100$, the second cluster of size $n_2 = 350$ is from the uniform distribution on a large “L”-shaped region, and the third cluster of size 50 is sampled from the uniform distribution on the entire $\mathbb{T}^2$. 
Data sets described above are generated for the estimation of prediction regions and clustering rules (displayed in Fig. 10). These data sets are called training data. Independent sets of data from the same models are also generated for validation, and are called testing data. Clustering rules based on the mixture models with $J, \alpha$ chosen by the proposed criterion (15) result in $K = 5$ clusters for Model I and $K = 2$ for Model II.

We have compared four methods of clustering: (i) the naive k-means clustering (ignoring the angular constraint), (ii) the extrinsic k-means clustering in the ambient space, and the proposed predictive clustering with membership assignment by (iii) the clustering rule $A_e$ (11) and by (iv) the clustering rule $A_o$ (12), creating an “outlier” cluster. See Sections 3.1 and 3.2 for details. For each model, the clustering rules are estimated using the training data, based on which the cluster memberships of the testing data are evaluated. Figure 11 displays the predicted cluster memberships for the testing data.

We now inspect the results for each model. Since Model I consists of five well-separated clusters, human eye is excellent in accurately clustering the data, with an understanding of the “cut-and-flattened” torus plots shown in Figs. 10–11. The naive k-means algorithm does not reflect the geometry of toroidal data, and results in a poor clustering (top left panel of Fig. 11). The extrinsic k-means correctly identifies the cluster #1 (split across $\phi = -\pi = \pi$), but mistakenly splits the elongated cluster (labeled 4 in Fig. 10) into two clusters (labeled 2 and 4 in Fig. 11). Our proposal correctly predicts all cluster memberships. To quantify the quality of clustering, we use the adjusted Rand index (Hubert and Arabie, 1985). For comparison of two cluster indices, one with $G$ clusters and the other with $K$ clusters, let $N_{gk}$ be the number of observations whose first cluster index is $g$ and the second index is $k$. The index is defined as

$$\text{ARI} = \frac{\sum_{g=1}^{G} \sum_{k=1}^{K} \binom{N_g}{2} - N_G N_K / \binom{N}{2}}{(N_G + N_K) / 2 - N_G N_K / \binom{N}{2}},$$

where $N_G = \sum_{g=1}^{G} \binom{N_g}{2}$ and $N_K = \sum_{k=1}^{K} \binom{N_k}{2}$. The ARI has the value of 1 when two indices match perfectly. Table 1 collects the ARIs comparing the predicted labels from each clustering method to the true cluster labels of the testing data. The proposed clustering with assignment rule $A_e$ performs almost perfectly for data from Model I.

The data from Model II can be viewed as two large clusters sprinkled with outliers (see right panels of Fig. 11). For this reason, we have used $K = 2$ in the k-means clusterings. As
before, the naive k-means algorithm is not suitable for toroidal data. The extrinsic k-means is not successful for this data set due to the irregularly shaped cluster. The proposed clustering methods correctly find the two large clusters. When the membership is assigned by $A_e$ (using the conformity score $\hat{c}_j(x)$), all observations belonging to the true “outlier” group are forced to be assigned to either the “L”-shaped cluster or the ball-shaped cluster. On the other hand,
the cluster membership rule $A_o$ with outliers performs better, as evidenced by the highest ARI in Table 1 and the visually satisfying clustering result in the lower right panel of Fig. 11.

5. Summary and discussion. We have introduced a clustering procedure for toroidal data based on the conformal prediction sets. Conformal prediction sets can be estimated using any conformity score, and we have demonstrated the use of the kernel density estimates, density estimates from a bivariate von Mises mixture model, and their variants in the construction of the conformity scores. The proposed clustering uses the mixture model-based prediction sets in identifying clusters and cluster membership assignment. We have shown that our approach performs better than naive adaptations of off-the-shelf clustering algorithms to toroidal data.

Our approach can be easily adapted for clustering the usual multivariate data in low-dimensional vector spaces. For data in the Euclidean space, the idea of using conformal prediction framework for clustering has also been explored by Shin, Rinaldo and Wasserman (2019) and Nouretdinov et al. (2020). Our proposal shares some similarity with these works but differs in the choices of conformal scores and the hyperparameters (as well as the different sample spaces). We also point out that the goal of Shin, Rinaldo and Wasserman (2019), building a prediction set based on clustering, is the opposite of this work: Cluster assignment based on prediction sets.

Our approach can be naturally extended to handle data on general tori $T^p (p \geq 2)$, directional data on hyperspheres $\{x \in \mathbb{R}^p : \|x\|_2 = 1\}$, or other types of manifold-valued data. We have focused on the bivariate circular variables $(\phi, \psi) \in T^2$, since the torsion angles have been known to well represent major backbone structures of proteins. For multivariate angles in a higher dimensional torus, fitting mixture models can be computationally expensive and unstable. A key direction for future research is to devise a conformity score that is computationally efficient for high $p$, and is easy to identify the resulting connected components (clusters). General tori $T^p$ appear as the data space for some biochemical applications. For example, the backbone of a protein has in fact a third angle, called $\omega$-angle, which are often ignored because $\omega \approx \pi$. If the protein has side chains, at each side chain there are a few torsion angles $\chi_i$ defined. In describing richer RNA structures, Sargsyan, Wright and Lim (2012) and Eltzner, Huckemann and Mardia (2018) have used seven torsion angles of RNA structures in $T^7$.

An important issue we have not fully investigated is the sampling distribution of $\hat{C}_n$. An empirical investigation in Appendix B.4 indicates that the variation due to $\hat{C}_n$ in $P(x \in \hat{C}_n | \hat{C}_n)$ is substantial. We leave this as a future topic of research.

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SUPPLEMENTARY MATERIAL

A. Algorithms. EM algorithms for mixtures of bivariate von Mises and an algorithm to test the intersection of two toroidal ellipses.
C. Supplementary figures. Supplementary figures referenced in Section 4.
D. Software. Data and R codes to reproduce the analysis are contained in a zipped file RcodesClustorus.zip. R functions used in the analysis are also available at https://github.com/sungkyujung/ClusTorus().

REFERENCES