# TORUS PRINCIPAL COMPONENT ANALYSIS WITH APPLICATIONS TO RNA STRUCTURE 

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#### Abstract

There are several cutting edge applications needing PCA methods for data on tori, and we propose a novel torus-PCA method that adaptively favors low-dimensional representations while preventing overfitting by a new test-both of which can be generally applied and address shortcomings in two previously proposed PCA methods. Unlike tangent space PCA, our torusPCA features structure fidelity by honoring the cyclic topology of the data space and, unlike geodesic PCA, produces nonwinding, nondense descriptors. These features are achieved by deforming tori into spheres with selfgluing and then using a variant of the recently developed principal nested spheres analysis. This PCA analysis involves a step of subsphere fitting, and we provide a new test to avoid overfitting. We validate our torus-PCA by application to an RNA benchmark data set. Further, using a larger RNA data set, torus-PCA recovers previously found structure, now globally at the onedimensional representation, which is not accessible via tangent space PCA.


1. Introduction. Dimension reduction on non-Euclidean manifolds with PCA-like methods has been a challenging task for which two usually successful categories of methods have been developed in the last decade: extrinsic (tangent space) approaches, for example, Arsigny et al. (2006), Boisvert et al. (2006), Fletcher et al. (2004), Gower (1975), and intrinsic (geodesic) ones, for example, Huckemann and Ziezold (2006). A critical review of PCA methods has been given in Huckemann, Hotz and Munk (2010); Sommer (2013) is another recently developed intrinsic PCA method. However, for the very simple non-Euclidean case of the flat and compact space of a torus (a direct product space of two or more angles), these approaches are not adequate. Namely, tangent space PCA (TS-PCA) fails to take into account the periodicity of the torus and, even worse, geodesic PCA is completely inapplicable because almost all geodesics densely wind around, as in Figure 1.

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(a) Flat torus as square in $\mathbb{R}^{2}$ with edges identified.

(b) Curved torus embedded in $\mathbb{R}^{3}$.

Fig. 1. Flat (a) and curved (b) torus representation. Except for horizontal and vertical geodesics (grey) in (a), and diagonal ones, all other geodesics wind around (a) and (b). All geodesics (black) with an irrational slope in (a) are dense.

In this paper we propose the novel tool of torus-PCA (T-PCA), which not only removes these defects but also more flexibly adapts to low dimension in a statistically controlled way to guard against overfitting. This is achieved by transforming the "geometrically benign" structure of the torus into a statistically benign geometry, namely one that does not allow for dense geodesics. We note that these dense geodesics are in the closure of the nondense geodesics, which in turn can be viewed as 1D subtori, and so an attempt for principal nested tori still suffers from the statistically nonbenign geometry. Specifically, we deform tori into spheres by choosing a nearby statistically benign geometry with clever mapping and self-gluing and then use a modification of the recently developed principal nested spheres analysis (PNS) of Jung, Dryden and Marron (2012). In particular, this PNS analysis involves a step of small sphere fitting and we provide a new test to avoid overfitting. However, deforming the geometry of the torus into that of a sphere, locally glued to itself (to honor periodicity), creates singularities (where curvature is unbounded). Notably, although locally respecting the flat geometry of the torus, ignoring periodicity, TS-PCA introduces in fact not only geometric but also topological singularities (the tangent space is not homeomorphic to the torus).

At this point we recall that within a sphere of radius $r>0$, every subsphere with the same radius $r$ is a great subsphere and one of smaller radius is a proper small subsphere. In this paper we speak of small subspheres to include great and proper small subspheres.

Some torus-specific PCA approaches have been developed apart from TS-PCA and geodesic PCA. Using wrapped normals, Kent and Mardia (2009) circumvent
the problem of winding geodesics and provide for an intrinsic parametric model with the same number of degrees of freedom as classical PCA. The PCA used by Altis et al. (2008) is a particular case of Kent and Mardia (2009). Allowing only geodesics that wind around at most once, as proposed by Kent and Mardia (2015), further reduces the degrees of freedom. As discussed in Huckemann and Eltzner (2015) for classical PCA in $\mathbb{R}^{n}$ the space of $k$-dimensional affine subspaces $(0 \leq k \leq n)$ has dimension $(n-k)(k+1)$; in contrast for PNS in the $n$ dimensional sphere, the space of $k$-dimensional small subspheres has dimension $(n-k)(k+2)(1 \leq k \leq n-1)$. For this reason (building on PNS), T-PCA more flexibly favors lower dimensional representations than TS-PCA, while this flexibility is better controlled against overfitting than in classical PNS.

Sargsyan, Wright and Lim (2012) may have been the first to treat toroidal data describing RNA structures in a spherical geometry. In their construction they halved the corresponding seven torus angles defined below and treated them as polar angles from a seven-dimensional sphere, thus taking only a very first step toward T-PCA. On this seven-dimensional sphere they investigated a test data set which we call the benchmark data. However, Sargsyan, Wright and Lim (2012) neither discussed nor exploited the drastic change of geometry, let alone amended by self-gluing, and only applied geodesic PCA [see Huckemann and Ziezold (2006)], maximizing projected variance and not minimizing residual variance. Incidentally, some pitfalls of using projected variance for compact manifolds were noted in Huckemann, Hotz and Munk (2010).

RNA structure analysis and challenges, a bigger picture. The last decades have witnessed finding an unexpected variety of RNA shape and function, and this variety is ever increasing. Base sequences, also called primary structures and consisting of polymers of four different nucleotides, are nowadays easily accessible by high throughput sequencing, and it is one ultimate goal to link these sequences to biological function. Biological function, however, is highly dependent on the 3D structure (or fold) which manifests at different levels [e.g., Brewer (2013), Chakrabarti, Chen and Varner (2011), Chapman, Sidrauski and Walter (1998), Seetin and Mathews (2012)]. At the bottom level is the single residue geometry usually described by dihedral angles between neighboring planes, each spanned by three adjacent atoms, similar to pages of an open book (Figure 2). The structure of each nucleotide can be described by six angles for the polymeric backbone and one angle for the nucleotide's base, giving a total of seven angles (Figure 3 and Table 1). Secondary structure is given by self-interaction within the RNA molecule via base pairing and other interactions forming specific patterns such as A-helices, hairpin loops and others. At the top level, tertiary and higher order structure arises from interacting lower order structure patterns via further base and backbone bindings.

In contrast to primary structure, the 3D structure is not easily accessible but needs to be reconstructed by elaborate technology such as X-ray crystallography.


FIG. 2. Illustration of a dihedral (torsion) angle defined by four atoms or three bonds, it is the opening angle between two pages of a book [reproduced from Mardia (2013)].

However, experimental structures are prone to misinterpretation and various errors. For example, backbone inconsistencies, where different reconstructed atoms occupy the same spatial location, frequently occur during reconstruction [Jain, Richardson and Richardson (2015), Richardson et al. (2008)]. To avoid or correct such errors, the space of possible 3D structures is often restrained or constrained to previously observed structures. This is typically done at the nucleotide or paired nucleotide level [Čech et al. (2013), Schneider, Morávek and Berman (2004), Wadley et al. (2007), Yang et al. (2003)]. Specifically, use is made of so-called rotamers describing empirical modes of probability distributions of nucleotide or nucleotide pair conformations. As these distributions are relatively peaked, limiting the conformational space to such rotamers avoids the introductions of incorrect conformations by limiting the conformational space to previously observed 3D patterns.

Among the many challenges along this path, we discuss two specific ones: data reduction methods and alignment strategies.

(a) $3 D$ structure of an $R N A$ residue.

(b) $2 D$ scheme of an $R N A$ residue.

FIG. 3. Part of an RNA backbone (phosphate groups with central atom denoted by P, followed by sugar rings that connect along the atoms labeled by $C 4^{\prime}$ and $C 3^{\prime}$, to which a nucleic base is bound). Dihedral angles (Greek letters) are defined by three bonds, the central bond carries the label; pseu-do-torsion angles (bold Greek letters) are defined by the pseudo-bonds between bold printed atoms [Figure 3(b)]. Underlying each pseudo-torsion angle are three heminucleotide angles. The precise definitions with same canonical atom notation are given in Table 1. The subscript "-" denotes angles of the neighboring residue. Figure 3(a) is reproduced from Frellsen et al. (2009).

TABLE 1
Atom bonds ( 2 nd column) defining angles ( 1 st column) with description ( 3 rd column). The two sets of heminucleotide angles (each of which can be approximated by a pseudo-torsion angle) define the backbone, which in conjunction with the base angle $\chi$ define a residue. Figure 3(a) shows the geometry of these atoms. ( $N$ denotes nitrogen)

```
\(\alpha O 3^{\prime}-P-O 5^{\prime}-C 5^{\prime}\)
\(\beta \quad P-O 5^{\prime}-C 5^{\prime}-C 4^{\prime}\) Heminucleotide angles
\(O 5^{\prime}-C 5^{\prime}-C 4^{\prime}-C 3^{\prime}\)
    \(C 5^{\prime}-C 4^{\prime}-C 3^{\prime}-O 3^{\prime}\)
    \(C 4^{\prime}-C 3^{\prime}-O 3^{\prime}-P \quad\) Heminucleotide angles
    \(C 3^{\prime}-O 3^{\prime}-P-O 5^{\prime}\)
    \(O 4^{\prime}-C 1^{\prime}-N 1-C 2\) For pyrimidine (monocyclic) bases
    \(O 4^{\prime}-C 1^{\prime}-N 9-C 4\) For purine (bicyclic) bases
    \(C 4^{\prime}-P-C 4^{\prime}-P\)
    \(P-C 4^{\prime}-P-C 4^{\prime}\)
    \(C 1^{\prime}-C 2^{\prime}-C 3^{\prime}-C 4^{\prime}\) Sugar pucker angle
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To the end of backbone reconstruction, single residue conformation space is explored and dimension reduction methods are applied to identify errors in experimental structures, provided among others by the popular free software of Davis et al. (2007). For example, removing inconsistencies, Murray et al. (2003) have found that RNA backbone is rotameric locally at hemi-nucleotide level, that is, among others, when reducing the 7D single residue space to a 3D backbone angular space, involving angles on only one side of the base (cf. Table 1 and Figure 3), conformer groups of each of the two sugar puckers (explained in Section 3) follow essentially one angle only. In our second application below, we revisit the data corresponding to one sugar pucker and generalize the result to finding a 1D structure common to all conformer groups.

On the one hand, matching RNA strands requires elaborate registration and alignment strategies [e.g., Mardia (2013)], building on statistical [e.g., Dryden and Mardia (2016), Srivastava and Klassen (2016)] and Bayesian [e.g., Green and Mardia (2006)] shape technology including non-Euclidean averaging and elastic curve representations [e.g., Laborde et al. (2013), Liu, Srivastava and Zhang (2011)]. On the other hand, averaging and exploring the 7D single residue space can be achieved via dynamically simulating similar structures [e.g., Chen and García (2013), Duarte and Pyle (1998), Estarellas et al. (2015)], and probabilistic approaches to this end require dimension reduction methods [e.g., Frellsen et al. (2009)]. In this context and also for higher-order structure prediction, it is necessary to explore not only the variation of single residue geometries typical for
specific secondary structure elements but also single residue geometries for intermediate and transition regions between structure elements [e.g., Dunbrack and Karplus (1994), Jain, Richardson and Richardson (2015)].

Applying torus-PCA to RNA structure analysis we provide for a novel dimension reduction method at residue level, and we apply it within the focus of current research to single residue geometries. However, it readily generalizes to simultaneous analysis of geometries of residue sequences ( $7 n$ angles for $n$ residues) but such an extension is left for future research. We measure effectively the statistical performance of our method by dimension reduction and faithfulness in terms of preserving previously known structure.

All of the angles used in our applications are defined in Table 1 and displayed in Figure 3. First we use the benchmark data set of Sargsyan, Wright and Lim (2012) which consists of neighborhoods of three known cluster centers in the $\eta-\theta$-plot [as in Figure 7(a), the pseudo-torsion angles $\eta, \theta$ are depicted in Figure 3(b); cf. also Table 1]. We find that T-PCA retrieves the underlying clusters in an effective way. This benchmark data set is a subset of a large RNA data set carefully selected for high experimental X-ray precision (0.3 nanometers) by Duarte and Pyle (1998), updated by Wadley et al. (2007) and analyzed by them and others, for example, Murray et al. (2003), Richardson et al. (2008). Next we use another subset of this large RNA data set with $\mathrm{C} 2^{\prime}$-endo sugar pucker (this and the other sugar pucker are explained fully in Section 3), subsequently called the C2 data set, where we compare our method to TS-PCA and show that T-PCA captures not only much more variance in the one-dimensional subspace but also the wrong topology in TS-PCA hides and tears apart subtle structural similarities.

In contrast, T-PCA provides structure fidelity, as global and local structural similarities are naturally preserved, most of it already visible in the 1D T-PCA representation, generalizing the above finding of Murray et al. (2003) that RNA backbone is locally rotameric at heminucleotide level, to:

These RNA conformers are rotameric at full residue level, possibly in a nonlinear sense, however.

In Section 2 we introduce torus PCA, which is the center-piece of our methodology. In Section 3 we apply our method to the benchmark and C2 data sets and review the results. The paper ends with a discussion and further illustrations in Supplement A [Eltzner, Huckemann and Mardia (2018a)]. An implementation of our T-PCA method and the RNA data sets we use are included as supplementary material, that is, Supplement B [Eltzner, Huckemann and Mardia (2018b)] and Supplement C [Eltzner, Huckemann and Mardia (2018c)], and can be found at http://www.stochastik.math.uni-goettingen.de/SFB755_B8.

Residues and residual variance. To avoid confusion, we clarify that the biochemical term residue denotes a RNA molecule segment corresponding to a single nucleic base (Section 3) whereas the statistical term residual variance denotes unexplained variation (Section 2.3).


FIG. 4. Self-gluing of $T^{2}$ : From a donut to a sausage. These operations are only topological; Figure 5 reflects the changes in geometry.
2. Torus PCA. Our dimension reduction procedure proceeds in two steps. First, the data space is deformed from a torus to a sphere with self-gluing, that is, parts of the sphere are topologically identified with themselves; see Figures 4 and 5. Several degrees of freedom are present in the deformation map we propose, and we discuss consequences of specific parameter choices. The second step is the dimension reduction for which we use a well-established procedure for dimension reduction on spheres with some extensions to take into account the original torus geometry and the self-gluing of the sphere.
2.1. Torus deformation schemes. Let $T^{D}=\left(\mathbb{S}^{1}\right)^{\times D}$ be the $D$-dimensional unit torus and $\mathbb{S}^{D}=\left\{x \in \mathbb{R}^{D+1}:\|x\|=1\right\}$ the $D$-dimensional unit sphere, $D \in \mathbb{N}$. The definition of the data-adaptive deformation mapping $P: T^{D} \longrightarrow \mathbb{S}^{D}$ defined in this section is based on comparing squared Riemannian line elements. If $\psi_{k} \in \mathbb{S}^{1}=[0,2 \pi] / \sim(k=1, \ldots, D)$ where $\sim$ denotes the usual identification of 0 with $2 \pi$, the squared line element of $T^{D}$ is given by the squared Euclidean line element

$$
d s_{T^{D}}^{2}=\sum_{k=1}^{D} d \psi_{k}^{2}
$$

For $\mathbb{S}^{D}$, in polar coordinates $\phi_{k} \in[0, \pi]$ for $k=1, \ldots, D-1$ and $\phi_{D} \in[0,2 \pi] / \sim$, whose relation to embedding Euclidean coordinates $x_{k}$ is given by

$$
\begin{aligned}
x_{1} & =\cos \phi_{1}, \\
\forall 2 \leq k \leq D: x_{k} & =\left(\prod_{j=1}^{k-1} \sin \alpha_{j}\right) \cos \phi_{k} \\
x_{D+1} & =\left(\prod_{j=1}^{D} \sin \phi_{j}\right)
\end{aligned}
$$



FIG. 5. All possibilities for gluing for $T^{2}$, illustrated by a data set uniform in a square with three outliers. Using mean centered (b), the square is near the equatorial great circle $\left[\psi_{1}=\pi\right.$ for $S O$ (e) and $\psi_{2}=\pi$ for $\left.S I(\mathrm{f})\right]$, and thus the square suffers little distortion, in comparison to the outliers. For gap centered (a), the outliers are less distorted, and for $S O$ (c) the square is particularly distorted because the equatorial great circle $\left(\psi_{1}=\pi\right)$ is then between outliers and square. In both cases, SO decreases the spread of the outliers, SI increases it, more drastically for mean centered. Due to the torus' periodicity, lines of same type in the flat torus angle plots [top row, (a) and (b)] are identified. The respective outer angle is halved, the respective inner angle is unscaled [middle row, (c) and (d)]. Due to collapsing of some identified lines to points (the singularity set, in Figure 4 this is the circle along which the donut is cut), north and south pole of each sphere are identified [bottom row, (e) and (f)].
the spherical squared line element is given by

$$
\begin{equation*}
d s_{\mathbb{S}^{D}}^{2}=d \phi_{1}^{2}+\sum_{k=2}^{D}\left(\prod_{j=1}^{k-1} \sin ^{2} \phi_{j}\right) d \phi_{k}^{2} \tag{1}
\end{equation*}
$$

In fact, this squared line element is not defined for the full sphere but only for $\phi_{k} \in(0, \pi)(k=1, \ldots, D-1)$, that is, the singularities of $\phi_{k}=0, \pi$ are excluded. The singularities at $\phi_{k}=0, \pi$ will account for singularities of $P$ which results in a self-gluing as explained below.

Angular distortions in a spherical geometry. Following colloquial usage, we use "distortion" as synonymous with "deformation" in the following. Because in (1), $d \phi_{1}^{2}$ comes with the factor 1 , no deformation at all occurs for $\phi_{1}$, that is, this angle corresponds to spherical distances without distortion. In the summation for $k=2$, we have a factor $\sin ^{2} \phi_{1}$ of $d \phi_{2}^{2}$, which shows how the angle $\phi_{1}$ distorts the angle $\phi_{2}$, and finally the deformation factor $\prod_{j=1}^{D-1} \sin ^{2} \phi_{j}$ of $d \phi_{D}^{2}$ reflects the distortions of $\phi_{D}$ by all other angles. For this reason, in the following we will refer to $\phi_{D}$ as the innermost angle and to $\phi_{1}$ as the outermost angle.

We now make an important note for later use:
REMARK 2.1. Near the equatorial great circle given by $\phi_{k}=\frac{\pi}{2} \quad(k=$ $1, \ldots, D-1$ ) the squared line element $d s^{2}$ is nearly Euclidean. Distortions occur whenever leaving the equatorial great circle. More precisely, distortions are higher when angles $\phi_{k}$ with low values of the index $k$ (outer angles) are close to zero or $\pi$, than when angles $\phi_{k}$ with high values of the index $k$ (inner angles) are close to zero or $\pi$.

DEFINITION 2.2 (Torus to Sphere Deformation). With a data-driven permutation $p$ of $\{1, \ldots, D\}$, data-driven central angles $\mu_{k}(k=1, \ldots, D)$ and data-driven scalings $\alpha_{k}$, all of which are described below, set

$$
\begin{equation*}
\phi_{k}=\frac{\pi}{2}+\alpha_{p(k)}\left(\psi_{p(k)}-\mu_{p(k)}\right), \quad k=1, \ldots, D \tag{2}
\end{equation*}
$$

where $p(k)$ is the index $k$ permuted by $p$ and the difference $\left(\psi_{p(k)}-\mu_{p(k)}\right)$ is taken modulo $2 \pi$ such that it is in the range $(-\pi, \pi]$.

We now explain in detail how the choices are data driven. Further illustration including practical advice is given in Supplement A [Eltzner, Huckemann and Mardia (2018a)]. First, we comment on the general applicability of T-PCA.

REMARK 2.3. The singularity set introduced, forms a subtorus of dimension $D-2$. In consequence, T-PCA is applicable whenever there is a structural data gap in all angles except for at most two-the larger the gap, the higher the structural fidelity.

In general, the scalings are restricted to the choices $\alpha_{k^{\prime}}=1 / 2$ and $\alpha_{k^{\prime}}=1$, $k^{\prime}=p(k)$. If all of the $k^{\prime}$ th torus angles of the data are within an interval of length $\pi$, choose $\alpha_{k^{\prime}}=1\left(k^{\prime}=1, \ldots, D-1\right)$ leading to unscaled $(\mathrm{U})$ angles. Otherwise, we choose $\alpha_{k^{\prime}}=1 / 2\left(k^{\prime}=1, \ldots, D-1\right)$ leading to halved $(\mathrm{H})$ angles. In practical
situations the torus data are often spread out over more than half circles for several angles. Then we choose (H) angles. In fact, for all of the analyses below, we chose $(\mathrm{H})$ angles and discuss below only the gluing effects corresponding to $(\mathrm{H})$ angles. Notably, the innermost angle $\phi_{D}$ always remains unscaled, $\alpha_{D}=1$. This is depicted in the second row of Figure 5.

The central angles $\mu_{k}$ will be chosen such that the mapped data points come to lie near the equatorial great circle and omit the singularities. Two plausible choices are:
(i) with the circular intrinsic mean $\bar{\psi}_{k \text {,intr }}$, set $\mu_{k}=\bar{\psi}_{k \text {,intr }}$ to obtain mean centered data;
(ii) with $\psi_{k, \text { gap }}$, the center of the largest gap between neighboring $\psi_{k}$ values of data points and $\psi_{k, \text { gap }}^{*}$ its antipodal point, define $\mu_{k}=\psi_{k, \text { gap }}^{*}$ to obtain gap (antipode) centered data.

While the implementation for (ii) is straightforward, for (i) we have used the fast algorithm from Hotz and Huckemann (2014). Mean centered data has the merit that the intrinsic means for each angle $\phi_{k}$ are mapped to the equatorial great circle thus minimizing deformation of the data.

For a strongly skewed data distribution, say spread out over a half circle, mean centered data using halved angles may touch the singularities, leading to high distortion there, while gap centered data will still be confined to a $\pi / 2$ neighborhood of the equator. On the other hand, for data sets with outliers, gap centered centering may be less robust than mean centered, making the latter more favorable, as depicted in Figures 5(c) and 5(e).

REMARK 2.4. Robustness w.r.t. outliers is surprisingly different on a compact space than on the usually considered noncompact spaces. Specific loci of outliers occurring nearly antipodal to the data bulk do not much affect the location of the mean, the largest data gap, however, is much more sensitive to these loci.

The choice of the permutation $p_{k}$ is driven by analyses of the data spread

$$
\begin{equation*}
\sigma_{k}^{2}=\sum_{i=1}^{n}\left(\psi_{k, i}-\mu_{k}\right)^{2}, \quad k=1, \ldots, D \tag{3}
\end{equation*}
$$

for each angle, where $\psi_{k, i} \in \mathbb{S}^{1}$ are the torus data and $n$ is the number of data points on $T^{D}$. If the angles are ordered by increasing data spread, such that $\sigma_{p(1)}^{2}$ is minimal and $\sigma_{p(D)}^{2}$ is maximal, in view of Remark 2.1, the change of distances between data points caused by the deformation factors $\sin ^{2} \phi_{j}$ in equation (1) is minimized. We call this ordering spread inside (SI), because variation is concentrated on the inner angles of the sphere. The opposite ordering is called spread outside (SO). Figure 5 illustrates different effects of SI and SO ordering of angles. We will restrict our considerations to these two options.

Self-gluing in case of halved angles, "From a donut to a sausage." In the following we give a brief overview of this procedure for $(\mathrm{H})$ halved angles [not for (U) angles for the reasons given above].

Due to periodicity on the torus, $\psi_{k}=0$ is identified with $\psi_{k}=2 \pi$ for all $k=1, \ldots, D$. In contrast, for all angles $\phi_{k}(k=1, \ldots, D-1), \phi_{k}=0$ denotes spherical locations different from $\phi_{k}=\pi$. For a representation respecting the torus' topology, however, it is necessary to identify these locations accordingly. Due to the spherical geometry, each of those regions is of dimension $D-j-1$, in which all angles vary except for $j$ of the $\phi_{1}, \ldots, \phi_{D-1}$ which are set to fixed values in $\{0, \pi\}$. In the topology of the torus, all those regions with a specific choice of fixed angles are identified with one another. In particular, there are $2(D-1)$ such regions of highest dimension $D-2$ on the sphere (where only one angle is fixed to 0 or $\pi$ ), two of which are pairwise identified in the topology of the torus. In fact, in the topology of the torus, each of these $D-1$ regions of highest dimension $D-2$ itself carries the topology of a torus of dimension $D-2$, each glued to each others' torus along a subtorus of dimension $D-3$ and so on. Thus the self-gluing of $S^{D}$ giving the topology of $T^{D}$ can be iteratively achieved along a topological subsphere of dimension $D-2$, which is suitably divided into $2(D-1)$ regions that are pairwise identified by way of a torus, sharing common boundaries which correspond to lower dimensional tori.

Example 2.5 details the case $D=3$, and Figures 4 and 5 illustrate the case $D=2$ as well as different choices for the permutation $p$.

Example 2.5. For $D=3$, on $\mathbb{S}^{3}$ we have the squared line element

$$
d s^{2}=d \phi_{1}^{2}+\sin ^{2} \phi_{1}\left(d \phi_{2}^{2}+\sin ^{2} \phi_{2} d \phi_{3}^{2}\right)
$$

where the angle ranges are $\phi_{1}, \phi_{2} \in[0, \pi], \phi_{3} \in[0,2 \pi)$.
Due to the spherical geometry in the region determined by $\phi_{1}=0 \bmod \pi$ or $\phi_{2}=0 \bmod \pi$, the circle $\phi_{3} \in[0,2 \pi)$ is a single point, say, $\phi_{3}=0$. This region is a topological circle on $S^{3}$ comprising four arcs

$$
\begin{array}{ll}
A_{1}=\left\{\left(0, \phi_{2}, 0\right): 0 \leq \phi_{2}<\pi\right\}, & A_{2}=\left\{\left(\pi, \phi_{2}, 0\right): 0 \leq \phi_{2}<\pi\right\}, \\
A_{3}=\left\{\left(\phi_{1}, 0,0\right): 0 \leq \phi_{1}<\pi\right\}, & A_{4}=\left\{\left(\phi_{1}, \pi, 0\right): 0 \leq \phi_{1}<\pi\right\} .
\end{array}
$$

Imposing the topology of the torus, when using halved angles, for $\phi_{1}$ and $\phi_{2}$ we also have the identification $0 \equiv \pi$ which results in the identification of $A_{1}$ with $A_{2}$ and of $A_{3}$ with $A_{4}$ with endpoints identified as one single point, forming a topological figure eight.
2.2. Linking the torus' deformation to PNS. For data sets on a torus, having applied a deformation on the resulting self-glued $\mathbb{S}^{D}$ (see Section 2.1 ), we modify principal nested sphere analysis (PNS) by Jung, Dryden and Marron (2012), Jung et al. (2010) for dimension reduction.

Assume a $d$-dimensional sphere $S^{d} \subset \mathbb{R}^{D+1}$ with center $x \in \mathbb{R}^{D+1}$ and radius $r>0$, and an affine $d$-dimensional plane $A^{d} \subset \mathbb{R}^{D+1}$ with distance $s<r$ from $x$. For $d \geq 2$ then the intersection $S^{d} \cap A^{d} \subset \mathbb{R}^{D+1}$ is a ( $d-1$ )-dimensional subsphere $S^{d-1}$ of $S^{d}$ with radius $r=\sqrt{1-s^{2}}$. If $r=1$ (i.e., $s=0$ ) this subsphere is a great subsphere, otherwise it is a proper small subsphere. For $d=1$ we pick just one point $\mu$, writing in expedient abuse of notation: $S^{0}=\{\mu\}$. In order to include all, great, proper small subspheres and the ultimate point, we call these small subspheres.

The PNS iteration leads to a sequence of small subspheres

$$
\begin{equation*}
\mathbb{S}^{D} \supset S^{D-1} \supset \cdots \supset S^{2} \supset S^{1} \supset S^{0}=\{\mu\} \tag{4}
\end{equation*}
$$

where the ultimate point $\mu$ is called the nested mean. Each $S^{d}(d=1, \ldots, D)$ is a $d$-dimensional sphere, the radii of which decrease monotonically with decreasing dimension (due to nesting). At each reduction step, the residual variances not explained by the corresponding subsphere are given as signed distances. Points lying inside the small subsphere-if it is a proper small sphere-receive a positive distance, points lying outside a negative distance. Indeed, for most realistic data applications, with probability one, all subspheres are proper small subspheres. However, to avoid overfitting, we want to ensure that the "small subsphere" is not too small but rather a great subsphere is fitted; see Section 2.4. In this case the direction of positive distance is picked at random. Similarly, we pick the direction of positive distance at random for the reduction from $d=1$ to $d=0$.

The classical PNS algorithm consists of two parts which alternate, namely the fitting of a subsphere $S^{d}$ and the projection to this subsphere $\pi_{d}: S^{d+1} \rightarrow S^{d}$ ( $d=D-1, \ldots, 0$ ) giving the fitted values explained by this subsphere. As $\mathbb{S}^{D}$ is glued to itself in T-PCA, distances through the glued part can be shorter than spherical distances. In such cases these distances are used in the fitting step as well as in the projection step. More precisely, our fitting procedure is done in two steps to avoid local minima. In the first step, we minimize the sum of squares of spherical distances. The resulting subsphere is taken as a starting point for the second step.

For the second step, we use the torus metric

$$
\delta: T^{D} \times T^{D} \rightarrow \mathbb{R}^{\geq 0} \quad(p, q) \mapsto\left(\sum_{i=1}^{D} \min \left(\left|p_{i}-q_{i}\right|^{2},\left(2 \pi-\left|p_{i}-q_{i}\right|\right)^{2}\right)\right)^{\frac{1}{2}}
$$

Assuming a data set $\mathcal{A}$ and a corresponding adaptive deformation $P_{\mathcal{A}}: T^{D} \rightarrow \mathbb{S}^{D}$ we define the following function on the sphere

$$
\begin{equation*}
\tilde{\delta}: \mathbb{S}^{D} \times \mathbb{S}^{D} \rightarrow \mathbb{R}^{\geq 0} \quad(x, y) \mapsto \delta\left(P_{\mathcal{A}}^{-1}(x), P_{\mathcal{A}}^{-1}(y)\right) \tag{5}
\end{equation*}
$$

using the inverse deformation $P_{\mathcal{A}}^{-1}$, which is well defined except for the singularities which are of dimension $D-2$. This is a metric when we take into account the topological identifications. To considerably lower computational speed for data
analyses, we orthogonally project data to lower dimensional subspheres using the spherical geometry only. On the deformed torus this can be viewed as a nonorthogonal projection. For the minimization in the second step, however, we use $\tilde{\delta}$ as the distance function.
2.3. Comparing variances. In Euclidean spaces, PCA variances are additive with monotone decrements leading to a convex variance plot as a property of the metric because decrements correspond to the nonincreasingly ordered eigenvalues of the corresponding covariance matrix. This means that every component can be thought of as contributing a fixed amount of variance and thus the sum of such individual variances can be understood as explained variance. If one views the principal components as defining a nested sequence of subspaces, the amount of variance which is not explained by the components spanning the subspace is equal to the residual variance of data around the subspace. Explained variance and residual variance add to 1 and thus yield equivalent descriptions of data variance.

In non-Euclidean spaces, linear PCA is not applicable and nonlinear dimension reduction methods do not come with a similar notion of additive variance [see the discussion for various definitions of intrinsic variances in Huckemann, Hotz and Munk (2010)]. This means that explained variance can no longer be defined in a straightforward way. However, residual variance is still a well-defined notion; therefore we use residual variances in the following to define cumulative variances and to compare results of different approaches.

Recall that T-PCA just as PNS yields a sequence of subspaces $\mathbb{S}^{D} \supset S^{D-1} \supset$ $\cdots \supset S^{1} \supset S^{0}=\{\mu\}$ with projections $\pi_{d}: S^{d+1} \rightarrow S^{d} \subset S^{d+1}(d=0, \ldots, D-1)$. From these we define the iterated projections

$$
\Pi_{d}=\pi_{d} \circ \pi_{d+1} \circ \cdots \circ \pi_{D-1}
$$

and finally the residual variances (variance not explained by $S^{d}$ ) of a data set $\mathcal{A}$

$$
V_{\mathcal{A}, P_{\mathcal{A}}, d}=\sum_{q \in \mathcal{A}} \tilde{\delta}^{2}\left(q, \Pi_{d}(q)\right), \quad d=0, \ldots, D-1
$$

and $V_{\mathcal{A}, P_{\mathcal{A}}, D}=0$, where $\tilde{\delta}$ is from (5). Due to nestedness, these sequences are nonincreasing with $d$. However, the decrements $V_{\mathcal{A}, P_{\mathcal{A}}, d-1}-V_{\mathcal{A}, P_{\mathcal{A}}, d}(d=1, \ldots, D)$ are not necessarily nonincreasing, so the resulting curve in the variance plot need not be convex. Still, this allows to define that $\{\mu\}, S^{1}, \ldots, S^{d}$ explain the cumulative variance up to dimension $d$

$$
V_{\mathcal{A}, P_{\mathcal{A}}, 0}-V_{\mathcal{A}, P_{\mathcal{A}}, d}, \quad d=0, \ldots, D
$$

which is nondecreasing in $d$.
2.4. Avoiding overfitting. In the PNS algorithm a cluster of points concentrated around a single center may still be best fitted by a "very" small subsphere. As this overfitting is obviously undesirable, Jung, Dryden and Marron (2012), Jung, Foskey and Marron (2011) have fitted a great subsphere in such cases; Jung, Foskey and Marron (2011) have given a decision rule whereas Jung, Dryden and Marron (2012) have given a test for this purpose. We propose the following new test based on a geometrically better suited model and highlight its attractive properties. In particular we show how robust our test is under the null model of Jung, Dryden and Marron (2012), which is a misspecified model for our case. We also indicate some limitations of the two previous procedures.

New model. Let $S^{d}$ be a fitted small subsphere, $2 \leq d<D$. For ease of notation, we now move and rescale $S^{d}$ to the unit sphere $\mathbb{S}^{d}$, without loss of generality, and $p \in \mathbb{S}^{d}$ is the center of the also moved and rescaled fitted small subsphere $S^{d-1} \subset$ $\mathbb{S}^{d}$. For our purpose, we can restrict our probability model for $q \in \mathbb{S}^{d}$, say, $g(q ; p)$, to depend only on the angular distance $r=d(p, q) \in[0, \pi]$. Further, suppose that vol $_{\mathbb{S}^{d}}$ denotes the surface volume of the $d$-dimensional unit sphere. Then, due to symmetry, $g$ fully characterizes the spherical angular marginal density of $r$

$$
\begin{equation*}
h(r ; p):=\operatorname{vol}_{\mathbb{S}^{d-1}} \cdot g(\gamma(r) ; p), \quad r \in[0, \pi] . \tag{6}
\end{equation*}
$$

Here, $\gamma$ is any curve along a great circle connecting $p$ with its antipodal, parametrized by $r \in[0, \pi]$ such that $\forall r: d(p, \gamma(r))=r$. Using the spherical volume element $d_{\mathbb{S} d} \Omega(q)$ at $q=\gamma(r)$ we note that

$$
1=\int g(q ; p) d_{\mathbb{S}^{d}} \Omega(q)=\int \frac{h(r ; p)}{\operatorname{vol}_{\mathbb{S}^{d}-1}} d_{\mathbb{S}_{d}} \Omega(q)=\int_{0}^{\pi} h(r ; p) \sin ^{d-1}(r) d r
$$

which means that $h(\cdot ; p)$ is indeed a marginal density with respect to the spherical angular measure

$$
d \mu(r)=\sin ^{d-1}(r) d r, \quad r \in[0, \pi]
$$

Then the Lebesgue angular marginal density $f(\cdot ; p)$ of $r$ is defined as

$$
f(r ; p):=\sin ^{d-1}(r) h(r ; p), \quad \int_{0}^{\pi} f(r ; p) d r=1
$$

since it gives the marginal density corresponding to $h(\cdot ; p)$ with respect to the Lebesgue measure on $[0, \pi]$.

Note that these densities are well studied for $d=2$ where the angle $r$ is called colatitude [see, e.g., Mardia and Jupp (2000)]; for the uniform distribution in polar coordinates for any $d$ on which this discussion is based, see, for example, Mardia, Kent and Bibby (1979).

For the following, we will need the density of the "folded normal distribution" on $[0, \infty)$ :

$$
\mathcal{F}(r ; \rho, \sigma):=\frac{1}{\sqrt{2 \pi} \sigma}\left(\exp \left(-\frac{(r-\rho \sigma)^{2}}{2 \sigma^{2}}\right)+\exp \left(-\frac{(r+\rho \sigma)^{2}}{2 \sigma^{2}}\right)\right)
$$

That is, we have

$$
\begin{equation*}
\mathcal{F}(r ; \rho, \sigma)=\frac{2}{\sqrt{2 \pi} \sigma} \exp \left(-\frac{r^{2}}{2 \sigma^{2}}-\frac{\rho^{2}}{2}\right) \cosh \left(\frac{r \rho}{\sigma}\right), \quad r \geq 0 \tag{7}
\end{equation*}
$$

This density has two positive parameters, $\rho$ and $\sigma$. Note that here $r$ is on $[0, \infty)$ so it is not restricted to $[0, \pi]$, a fact which will be of importance later on where we will truncate this distribution. For $\rho \rightarrow \infty$ this tends to a usual normal distribution centered at $\rho \sigma$, while it becomes a halved normal distribution (of doubled height) for $\rho \rightarrow 0$. For $\rho \leq 1$ the mode stays fixed at the origin; for $\rho>1$ it moves to the right.

With the above discussion on the marginals we therefore choose $g \propto \mathcal{F}$ yielding the spherical angular marginal density $h$ and the Lebesgue angular marginal density $f$ :

$$
\begin{align*}
& h(r ; p, \rho, \sigma):=\frac{\sqrt{2 \pi} \sigma}{\mathcal{C}(\rho, \sigma)} \mathcal{F}(r ; \rho, \sigma), \\
& f(r ; p, \rho, \sigma):=\frac{\sqrt{2 \pi} \sigma}{\mathcal{C}(\rho, \sigma)} \sin ^{d-1}(r) \mathcal{F}(r ; \rho, \sigma), \quad r \in[0, \pi] \tag{8}
\end{align*}
$$

where we have truncated $\mathcal{F}(r ; \rho, \sigma)$ from (7) and $\mathcal{C}(\rho, \sigma)$ is the normalization. These will be referred to as $h$ - and $f$-distribution, respectively, in the following.

Subsequently, it will be important to note the following property of these distributions, for dimension $d=2$, as a surface of revolution over $\mathbb{R}^{2}$. In polar coordinates $(r, \vartheta) \mapsto \mathcal{F}(r ; \rho, \sigma) \frac{1}{2 \pi}$, the case $\rho>1$ yields a ring while the case $\rho=0$ yields a symmetric Gaussian distribution. Due to its smoothness it is a good candidate for a test distribution for the angular spherical marginal density (6) to distinguish "just" concentrated data near $p$ ( $p$ is at $r=0$ ) from concentrated data along a distinct subsphere (a ring in 2D) around $p$.

Likelihood ratio test. Suppose we are given the sample $\left\{q_{1}, \ldots, q_{n}\right\}$ from the $f$-distribution with the spherical distances $r_{i}=d\left(p, q_{i}\right)(i=1, \ldots, n)$ where the center $p$ of the subsphere is known. If $\rho \leq 1$, the $h$ distribution has its maximum at $r=0$, that is, there is no proper small spherical structure about the center $p$. If $\rho>1$, there is a proper small spherical structure about the center $p$. Thus, $\rho=1$ forms the boundary between the two cases.

Therefore, we can formulate our hypotheses as follows for testing for a great subsphere:

$$
\begin{equation*}
H_{0}: \rho=1 \text { (great subsphere) } \quad \text { vs. } H_{1}: \rho>1 \text { (small subsphere). } \tag{9}
\end{equation*}
$$

The log likelihood up to a constant is given by

$$
\begin{aligned}
\ell\left(\rho, \sigma \mid\left\{r_{i}\right\}_{i=1}^{n}\right)= & -n \ln \mathcal{C}(\rho, \sigma)+(d-1) \sum_{i=1}^{n} \ln \sin \left(r_{i}\right) \\
& -\frac{n \rho^{2}}{2}-n \ln (\sigma)+\sum_{i=1}^{n}\left(-\frac{r_{i}^{2}}{2 \sigma^{2}}+\ln \cosh \left(\frac{r_{i} \rho}{\sigma}\right)\right) .
\end{aligned}
$$

Note that the normalization $\mathcal{C}(\rho, \sigma)$ can be easily computed numerically so we can determine the MLEs for $\rho$ and $\sigma$ using standard numerical optimization. For $H_{1}$, the MLEs need to be constrained under $\rho>1$. Then twice the log of the likelihood ratio (with negative sign) is given by

$$
\begin{align*}
\lambda= & 2 \sup \left\{\ell\left(\rho, \sigma \mid\left\{r_{i}\right\}_{i=1}^{n}\right): \rho \in(1, \infty), \sigma \in \mathbb{R}^{+}\right\}  \tag{10}\\
& -2 \sup \left\{\ell\left(\rho, \sigma \mid\left\{r_{i}\right\}_{i=1}^{n}\right): \rho=1, \sigma \in \mathbb{R}^{+}\right\} .
\end{align*}
$$

From Wilks' theorem, the statistic $\lambda$, under $H_{0}$, is asymptotically distributed as $\chi_{1}^{2}$. We use a $5 \%$ significance level for our test, which means that when $H_{0}$ is rejected, we keep the fitted small subsphere if $\lambda>\chi_{1,0.95}^{2} \approx 3.84$; otherwise, we perform a great subsphere fit.

Comparison with the decision rule of Jung, Foskey and Marron (2011). This rule is based on another type of angular $h$ and $f$ [versus our angular $f$ and $h$ given by (8)]

$$
\begin{aligned}
h_{\mathrm{Jung}}(r ; p, \rho, \sigma) & :=\frac{1}{\sin ^{d-1}(r)} \mathcal{F}(r ; \rho, \sigma), \\
f_{\mathrm{Jung}}(r ; p, \rho, \sigma) & :=\mathcal{F}(r ; \rho, \sigma) .
\end{aligned}
$$

In their decision rule, a great sphere is fitted if the probability distribution does not exhibit a ring-shaped local maximum, which is the case if $\rho \leq 2$. But this model leads to a singularity of the density $h_{\text {Jung }}$ at $p$, which is not a desirable feature. In contrast, our $h$-distribution leads to a smooth distribution on the sphere as illustrated by the above considerations about the surface of revolution. Our $h$ distribution is compared with the $h_{\mathrm{Jung}}$ distribution in Figure 6 for appropriate values of $\rho$. Our $h$ distribution is the same for all $d$ but for illustration, we have used $d=2$ for $h_{\text {Jung }}$ which depends on $d$.

In validation of our test we carried out two simulation studies:
$D_{0}$ : We simulate data under $H_{0}$ in (9) by choosing $\rho=1$ in (8) and average over the nuisance parameter $\sigma$ by 1000 samples, where in each $\sigma$ is uniform in [0.1, 0.4].
$D_{1}$ : We simulate data under $H_{1}$ in (9) by choosing various combinations of $\rho \in$ $\{1.2,1.5,2,3\}$ and $\sigma \in\{0.15,0.2,0.5\}$ in (8), and for each we average using 1000 samples.

The results in Table 2 show that our test at asymptotic level of 5\%, that is, it rejects a small sphere when $\lambda>\chi_{1,0.95}^{2} \approx 3.84$ with $\lambda$ from (10), holds asymptotically the level and that the Type 2 error asymptotically decays to zero, very quickly for larger $\rho$. Since for $N=100,150$ and 200 the true levels are above $5 \%$, we have estimated the asymptotic levels yielding a true level of $5 \%$ in Table 3 and display the corresponding Type 2 error there also. This estimation is a matter of minutes for $N=100$ and below one hour for $N=1000$. Based on these simulations and as a rule of thumb, we recommend using our test for sample sizes


Fig. 6. The probability densities for $\sigma=0.5$ along the geodesic $\gamma$ in $\mathbb{S}^{d}$ from (6) for our $h$ (invariant under $d$ ) and the $h_{\text {Jung }}($ for $d=2$ ) distribution. Displaying a value for $\rho$ below the respective boundary, at the boundary and above the boundary; namely, $\rho=1$ for our $h$ and $\rho=2$ for the $h_{\text {Jung }}$ distribution.
beginning with $N=300$. This is certainly the case for our application to the C2 data set with $N=649$; cf. Section 3.2. On the benchmark data set with $N=181$, as discussed in Section 3.1, our test also does well, although the sample size seems slightly below the asymptotic regime. For both data sets we have used our test against overfitting a small sphere at asymptotic level of 5\%.

Assessment of robustness of our test under the null distribution of Jung, Dryden and Marron (2012). We now assess the robustness of our test under a misspecified model, namely, the von Mises-Fisher distribution, which is the null distribution of Jung, Dryden and Marron (2012). To carry this out, we note the following points related to their test. First, we note that they have translated their null hypothesis

TABLE 2
Type 1 errors (rejecting $H_{0}$ ) for $D_{0}$ and Type 2 errors (accepting $H_{0}$ ) for $D_{1}$ for our test with various parameter values in a simulation with 1000 repetitions and asymptotic level of $5 \%$, that is, rejecting for $\lambda>\chi_{1,0.95}^{2} \approx 3.84$ with $\lambda$ from (10)

| Sample size | Type $1\left(D_{0}\right)$$\rho=1$ | Type $2\left(D_{1}\right)$ |  |  |  |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  |  | $\begin{gathered} \rho=1.2 \\ \sigma=0.15 \end{gathered}$ | $\begin{aligned} & \rho=1.5 \\ & \sigma=0.2 \end{aligned}$ | $\begin{gathered} \rho=2 \\ \sigma=0.2 \end{gathered}$ | $\begin{gathered} \rho=2 \\ \sigma=0.5 \end{gathered}$ | $\begin{gathered} \rho=3 \\ \sigma=0.15 \end{gathered}$ |
| 100 | 7.4\% | 80.4\% | 41.2\% | 3.4\% | <0.1\% | <0.1\% |
| 150 | 6.1\% | 79.9\% | 29.3\% | 0.5\% | <0.1\% | $<0.1 \%$ |
| 200 | 5.5\% | 73.2\% | 20.2\% | <0.1\% | <0.1\% | $<0.1 \%$ |
| 300 | 5.0\% | 71.2\% | 8.3\% | <0.1\% | <0.1\% | $<0.1 \%$ |
| 500 | 5.0\% | 59.7\% | 1.0\% | <0.1\% | <0.1\% | $<0.1 \%$ |
| 1000 | 4.9\% | 34.7\% | <0.1\% | $<0.1 \%$ | $<0.1 \%$ | $<0.1 \%$ |

TABLE 3
We estimate the asymptotic level for our test leading to a true level of 5\%, that is, achieving a Type 1 error (rejecting $H_{0}$ ) for $D_{0}$ of $5 \%$. The table gives the asymptotic level and Type 2 errors (accepting $H_{0}$ ) for $D_{1}$ for our test with various parameter values in a simulation with 1000 repetitions

| Sample size | $\begin{gathered} \text { Asymptotic level } \\ \left(D_{0}\right) \\ \rho=1 \end{gathered}$ | Type $2\left(D_{1}\right)$ |  |  |  |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  |  | $\begin{gathered} \rho=1.2 \\ \sigma=0.15 \end{gathered}$ | $\begin{aligned} & \rho=1.5 \\ & \sigma=0.2 \end{aligned}$ | $\begin{gathered} \rho=2 \\ \sigma=0.2 \end{gathered}$ | $\begin{gathered} \rho=2 \\ \sigma=0.5 \end{gathered}$ | $\begin{gathered} \rho=3 \\ \sigma=0.15 \end{gathered}$ |
| 100 | 3.0\% | 85.0\% | 54.4\% | 5.1\% | <0.1\% | $<0.1 \%$ |
| 150 | 4.0\% | 81.1\% | 31.8\% | 0.9\% | <0.1\% | <0.1\% |
| 200 | 4.4\% | 76.4\% | 24.2\% | 0.1\% | $<0.1 \%$ | $<0.1 \%$ |
| 300 | 5.0\% | 71.2\% | 8.3\% | <0.1\% | $<0.1 \%$ | <0.1\% |
| 500 | 5.0\% | 59.7\% | 1.0\% | $<0.1 \%$ | $<0.1 \%$ | <0.1\% |
| 1000 | 5.0\% | 34.7\% | <0.1\% | <0.1\% | <0.1\% | $<0.1 \%$ |

of a compact cluster into fitting by a great subsphere through a von Mises-Fisher distribution. The parameters of this distribution are estimated via MLE. Then a Student $t$-like test statistic of distances to the estimated center point is used as their test statistic. Next, we note that for their test statistic, they simulate bootstrap quantiles from the von Mises-Fisher distribution with parameters given by the MLE. However, Jung, Dryden and Marron (2012) have given neither a theoretical result (like we have the asymptotic p-value of our test statistics $\lambda$ ), nor a simulation study to assess their test statistics under their null hypothesis. We have reimplemented their data driven procedure so as to use their null hypothesis and have carried out the following simulation study:
$D_{0}^{\prime}$ : Here, we directly simulate spherical samples leading to a great circle, from the null hypothesis of the test of Jung, Dryden and Marron (2012), namely from a von Mises-Fisher distribution with density in $x$ proportional to $e^{\kappa \mu^{T} x}$ with a high value of the concentration parameter $\kappa=10$ to give a fair chance. We average over 1000 samples with $\mu$ uniform on the sphere.

As shown in Table 4, we note that our test is more conservative on the null hypothesis of the test of Jung, Dryden and Marron (2012). Further, the true level of the test of Jung, Dryden and Marron (2012) also decreases with sample size, and almost reaches the simulated level for $N=1000$. In passing, we note that estimating the simulated level leading to a true level of $5 \%$ for the test by Jung, Dryden and Marron (2012), however, is impractical, as for $N=100$ already, estimation takes weeks.
3. Application to RNA structure. RNA is usually single stranded, and the single strand interacts with itself and forms complex shapes (this is in contrast to DNA which usually takes a double-stranded helical conformation). This means

TABLE 4
Type 1 errors [rejecting the null hypothesis of Jung, Dryden and Marron (2012) which is a von Mises-Fisher distribution] for the test of Jung, Dryden and Marron (2012) and errors under this misspecified model for our test, with concentration parameter $\kappa=10$ in a simulation with 1000 repetitions. For their test we use a simulated level of 5\% and for our test we use an asymptotic level of $5 \%$

| Sample size | Jung, Dryden and Marron (2012) | Our test |
| :--- | :---: | ---: |
| 100 | $17.0 \%$ | $1.0 \%$ |
| 150 | $13.4 \%$ | $0.2 \%$ |
| 200 | $13.4 \%$ | $<0.1 \%$ |
| 300 | $8.8 \%$ | $<0.1 \%$ |
| 500 | $8.4 \%$ | $<0.1 \%$ |
| 1000 | $5.9 \%$ | $<0.1 \%$ |

that the geometry is rather variable even on the scale of single atoms. As described in Section 1, each nucleic base corresponds to a backbone segment described by 6 dihedral angles and one angle for the base, giving a total of seven angles; cf. Table 1 and Figure 3. The distribution of these seven angles over large samples of RNA strands have been studied in detail, see Frellsen et al. (2009), Murray et al. (2003), Richardson et al. (2008), Schneider, Morávek and Berman (2004), Wadley et al. (2007). Figure 3(a) details a segment of the RNA backbone with seven angles for each residue giving the 3D folding structure. An approximation of the geometric folding structure on the level of single residues is given by the two pseudo-torsion angles $\eta$ and $\theta$ [Figure 3(b)]. These two (dihedral) angles provide at once a two-dimensional visualization [Figure 7(a)]; see, for example, Duarte and Pyle (1998), Wadley et al. (2007).

Finally, the dihedral angle $\nu_{2}$ [Figure 3(b) and Table 1] quantifies the folding (pucker) of the sugar ring. Only two modes of folding are geometrically and energetically possible, which are characterized by either $\mathrm{C} 3^{\prime}$ or $\mathrm{C} 2^{\prime}$ being outside the plane spanned by $\mathrm{C1}^{\prime}-\mathrm{O}^{\prime}-\mathrm{C} 4^{\prime}$ and toward the direction of $\mathrm{O}^{\prime}$. If $\mathrm{C} 2^{\prime}$ lies outside the plane then $\nu_{2} \approx 325^{\circ}$, this is called $C 2^{\prime}$-endo sugar pucker; whereas if $\mathrm{C} 3^{\prime}$ lies outside the plane then $\nu_{2} \approx 35^{\circ}$, this is called $C 3^{\prime}$-endo sugar pucker. The hydroxy group attached to the $\mathrm{C} 2^{\prime}$ atom in RNA causes the $\mathrm{C} 3^{\prime}$-endo sugar pucker to be energetically preferred [see, e.g., Egli, Portmann and Usman (1996)], and thus this is about 10 times more abundant than the $\mathrm{C}^{\prime}{ }^{\prime}$-endo sugar pucker in the large RNA data set of Duarte and Pyle (1998) and Wadley et al. (2007).

For our application below we use two subsets of a large classical data set (8301 residues), which was carefully selected for high experimental X-ray precision ( 0.3 nanometers) by Duarte and Pyle (1998), updated by Wadley et al. (2007) and analyzed by them and others, for example, Murray et al. (2003), Richardson et al. (2008).


Fig. 7. (a) The benchmark data set of Sargsyan, Wright and Lim (2012) with their three preselected clusters in the $\eta-\theta$ plot. (b) The benchmark data set plotted for the two most discriminant angles ( $\alpha$, $\zeta)$ chosen out of the seven dihedral angles; in the "donut to sausage" transformation along the dashed lines the corresponding angles are collapsed to a single point.
3.1. The benchmark data set. This benchmark data set has been carefully selected by Sargsyan, Wright and Lim (2012) to validate their method. From the C3'endo sugar pucker they took clusters labeled I ("triangles," 59 points), II ("crosses," 83 points) and V ("disks," 39 points) by Wadley et al. (2007) totaling 181 data points, which form three clusters in the $\eta-\theta$ plot as shown in Figure 7(a). While clusters I and II correspond to distinct structural elements featuring base stacking, the residues in cluster V belong to a wider variety of structural elements.

Visualization is obviously not possible in the 7D space of all torsion angles. However, we find that the angle pair $(\alpha, \zeta)$ is the most discriminatory, and a plot is given in Figure 7(b). The "disks" cluster is not very concentrated, in contrast to the "crosses" cluster which is twice as big. Parts of the "disks" are very close to the "crosses" cluster. In fact upon close inspection, due to periodicity, the "triangles" and "crosses" clusters are also rather close in the $\eta-\theta$ plot in Figure 7(a).

We have applied T-PCA to all seven angles and depict the two-dimensional representation for SI ordering in Figure 8(a) (which is hardly visually distinguishable from SO ordering). To see that the data are, in fact, very well approximated by the best fit circle we use a planar representation of the first two T-PCs in Figure 8(b). Using the same symbols for Figure 8 as in Figure 7 shows that the three preselected clusters can be rather well distinguished by eye. We note that the first component explains $84 \%$ of data variation. In comparison in Figure 8(c) we adapt Figure 6 from Sargsyan, Wright and Lim (2012). Again the clusters can be well discriminated along the first GeoPC [horizontal in the 2D approximation in Figure 8(b)]. In contrast to T-PCA, however, the data are not well approximated by the

(a) 2D approximation, T-PCA (SI)

(b) $2 D$ approximation, $T-P C A$ (SI) in planar coordinates

(c) 2D approximation, GeoPCA

Fig. 8. Two-dimensional PCA approximations of the benchmark data set via T-PCA with SI ordering in natural spherical coordinates (a), in planar coordinates (b) and GeoPCA adapted from Sargsyan, Wright and Lim (2012), Figure 6, (c). The symbols represent the same clusters as in Figure 7.
first GeoPC, as the projections to the second GeoPC component [vertical in the 2D approximation in Figure 8(c)] feature maximal data range. In fact, both GeoPCs explain roughly similar amounts of data variation.

Thus Figure 8 illustrates the power of T-PCA going significantly beyond the analysis of Sargsyan, Wright and Lim (2012). Not only can the preselected clusters be separated but the data are very accurately approximated by their projection to the 1D component.
3.2. The $1 D$ structure of $C 2$ data set. We now describe in detail how our C 2 data set is extracted from the large RNA data set. Notably, some of the RNA structures in this data set are only short pieces adhering to a protein or another RNA structure. Therefore, we prune by removing residues further than $50^{\circ}$ in torus distance from their nearest neighbor. This leads to 7,544 residues and 649 of these are residues with $\mathrm{C} 2^{\prime}$-endo sugar pucker, that is, $\nu_{2} \in\left[300^{\circ}, 350^{\circ}\right]$. This produces a moderately large data set to analyze (in contrast to the very large data set of all other residues including $\mathrm{C} 3^{\prime}$-endo sugar pucker).

Murray et al. (2003) noted that this data set is locally rotameric, as, among others, conformer clusters essentially extend along the $\beta$ angle, considering only the three heminucleotide angles $\alpha-\beta-\gamma$ [Figure 9(a)]. Already in this heminucleotide space, these individual 1D cluster patterns compete with the group spread along the $\alpha$ angle and in full 7D residual space; there are more competing features, which, in the 2D TS-PCA plot involving all seven angles, manifest as three diffused stripe shaped clusters [Figure 9(b)]. Here the 1D pattern of the largest conformer group can be traced along the shifted second diagonal. The two conformer groups next in size, which are close in heminucleotide angles, are ripped apart in TS-PCA due to its wrong topology, because they are far from the base point of the tangent space that is controlled by the dominating cluster. Notably, the correct topology could not even be forced onto that plot because, due to the winding effects illustrated in Figure 1, boundary loci correspond to different torus loci.

Due to its larger flexibility and higher fidelity, T-PCA recovers a 1D pattern as the overall dominating structure, reflecting the proximity of the second and third largest cluster in the second component [Figure 9(c) and (d) in planar representation for better illustration, which is, of course, periodic]. Notably, according to Remark 2.3, structural fidelity can be expected due to the large gaps in the $\beta$ and $\gamma$ angles; cf. Figure 9(a). Using T-PCA, we generalize the finding of a locally rotameric structure by Murray et al. (2003) to:

> In full 7D angular space, the RNA residue conformers are rotameric, essentially following a single angle that is a nonlinear combination of the original ones, however.

Upon yet closer inspection, the fine clustering along the 1 D component reflects the clustering in the complementary heminucleotide $\delta-\varepsilon-\zeta$ angles from Murray et al. (2003), Figure 4(c), rear part.
3.3. Comparing T-PCA with TS-PCA. We summarize our use of T-PCA and TS-PCA using all seven angles for the C2 data in Table 5(a) and Figure 10(a). In 1D, T-PCA captures $73 \%$ of the variance whereas TS-PCA captures only $44 \%$ of the variance. Only when adding a second dimension TS-PCA captures more variance ( $81 \%$ ) than the 1D component of T-PCA. Higher order PCs, both for TPCA and TS-PCA, explain roughly the same amount of data variance.

(a) Conformer clusters in the $\alpha-\beta-$ $\gamma$ plot adapted from (Murray et al., 2003, Figure 4.b)

(c) T-PCA (SI)

(b) TS-PCA

(d) T-PCA (SI), planar view

Fig. 9. Residues with C2'-endo sugar pucker with clustering following Murray et al. (2003). Three--dimensional heminucleotide angles (a); two-dimensional TS-PCA (b) approximation; two-dimensional T-PCA (SI) approximation, the small circle gives the 1D approximation (c); two-dimensional T-PCA (SI) approximation in planar representation (d).

To highlight the differences between the two PCA methods, let us consider the example of three points. There is an exactly fitting small circle used by T-PCA. Indeed, if applied to the $\eta-\theta$ plot [Figure 7(a)], T-PCA would reduce the three clusters rather accurately to a 1D circle. In contrast, TS-PCA approximates three points only along a straight line in the tangent space, and such an approximation is only possible if data lie favorably such as in the $\eta-\theta$ plot; see Figure 7(a). The

TABLE 5
Cumulative variances for T-PCA (SI) and for TS-PCA

|  | (a) C2 data |  |  |  | (b) Simulated simplex data |  |  |
| :--- | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Dimension | T-PCA (SI) | TS-PCA |  | Dimension | T-PCA (SI) | TS-PCA |  |
| 1 | $74 \%$ | $44 \%$ |  | 1 | $39 \%$ | $18 \%$ |  |
| 2 | $83 \%$ | $81 \%$ |  | 2 | $50 \%$ | $34 \%$ |  |
| 3 | $90 \%$ | $90 \%$ |  | 3 | $63 \%$ | $48 \%$ |  |
| 4 | $95 \%$ | $94 \%$ |  | 4 | $77 \%$ | $62 \%$ |  |
| 5 | $98 \%$ | $97 \%$ |  | 5 | $89 \%$ | $75 \%$ |  |
| 6 | $99 \%$ | $99 \%$ |  | 6 | $95 \%$ | $88 \%$ |  |

$\alpha-\zeta$ plot [Figure 7(b)], however, illustrates that a 1D approximation for all seven angles is not possible for TS-PCA, while it is possible for T-PCA [Figure 8(b)].

In fact, usually T-PCA requires one dimension less than TS-PCA because $k$ points in general position span a $k$-dimensional affine subspace, which is detected by TS-PCA, and the surface of a $(k-1)$-dimensional sphere, which is detected by T-PCA. We illustrate this using a simulated simplex data set with points in general position, namely, 8007 D angles distributed independently at one of eight simplex vertices, $\pi$ apart with Gaussian noise of variance $(\pi / 3)^{2}$. The results are displayed in Table 5(b) and Figure 10(b). If there are affine data dependencies, however, this advantage of T-PCA over TS-PCA by one dimension is lost. Indeed the C2 data set features such affine dependencies between angles, which is already visible in Figure 9(a), and hence in Figure 10(a), T-PCA outperforms TS-PCA in terms of explained variance only in dimension one.
4. Discussion. We have provided a novel framework for torus PCA to perform PCA-like dimension reduction for angular data. Previous attempts have not been


FIG. 10. Scree plots of cumulative variances for T-PCA (SI) compared to TS-PCA.
satisfactory, because, on the one hand, the geometry featuring dense geodesics leads to severe restrictions for geodesic approaches while, on the other hand, Euclidean approximations disregard periodicity. We have used an adaptive deformation to a statistically benign geometry, allowing for increased and statistically controlled flexibility whilst at the same time guaranteeing structure fidelity. In application to dihedral angles of RNA structures we have validated our method using a classical benchmark data set. Using a C2'-endo sugar pucker residue data set we have given evidence on how T-PCA is better and more meaningful than TS-PCA, and we have illustrated that the significant interdependence found by Murray et al. (2003) in a 3D representation is seen by T-PCA remarkably in 1D.

There are several benefits coming with dimension reduction to 1D. In view of data clustering, it allows to build on powerful and well established statistical 1D methods for mode detection [e.g., Dümbgen and Walther (2008), Huckemann et al. (2016), Schmidt-Hieber, Munk and Dümbgen (2013)], and this challenge will be taken up in future research.

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## SUPPLEMENTARY MATERIAL

Supplement A: Data (DOI: 10.1214/17-AOAS1115SUPPA; .pdf). An illustration how to choose data-driven parameters for torus PCA.

Supplement B: Data (DOI: 10.1214/17-AOAS1115SUPPB; .zip). RNA residue data used for the analysis in this paper.

Supplement C: Implementation (DOI: 10.1214/17-AOAS1115SUPPC; .zip). Source code of the T-PCA implementation used for this paper.

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