

Persistent Khovanov Homology for Knots and Banded Knots

Matthew Zabka

Department of Mathematics and Statistics, Southwest Minnesota State University

Email: matthew.zabka@smsu.edu

Anzor Beridze

School of Mathematics and Computer Science, Kutaisi International University

Email: anzor.beridze@kiu.edu.ge

Draft Proposal for Internal and Interdisciplinary Discussion

(This document is a preliminary working draft intended for conceptual development and collaborative review. Additional authors may be added as the project evolves.)

Introduction

Knot theory has emerged as a powerful framework for understanding the geometry and topology of biological macromolecules. DNA often becomes knotted or linked during replication, transcription, and packaging, and cells rely on specialized enzymes such as topoisomerases to regulate this topological complexity [1, 2]. Proteins, too, can display deeply knotted or slipknotted configurations whose folding pathways and functional implications remain the subject of active investigation [3]. At larger scales, chromatin and chromosomes exhibit intricate polymeric entanglements that influence their spatial organization and epigenetic state [4]. Even RNA, although generally thought to avoid deep knots, can exhibit nontrivial entanglements and topological features in three-dimensional structural models [5]. These examples illustrate that knot theory—including both classical knot invariants and newer homological tools—is well positioned to contribute to the quantitative characterization of complex, noisy, or partially observed biological structures.

Despite this widespread appearance of knotted and entangled structures in biology, in many experimental settings the topology of a macromolecule cannot be determined exactly. Cryo-electron microscopy, for example, often produces three-dimensional density maps with unresolved or ambiguous loop regions, leaving substantial uncertainty about the underlying geometric configuration [6]. Chromatin conformation capture techniques such as Hi-C provide only pairwise contact frequencies rather than full spatial embeddings, making it impossible to determine a definitive knot or link type from the data [7]. Even when explicit polymer models are used to reconstruct chromosome geometry, the resulting embeddings may display topological features—including knots, links, and slipknots—that fluctuate across model realizations or remain only partially localized [8, 9]. These challenges highlight the need for mathematical tools capable of quantifying topological structure in settings where only incomplete, noisy, or ensemble-based information is available. Persistent homology provides a natural framework for addressing such uncertainty, allowing knot-like features to be detected, tracked, and compared across varying spatial scales or model parameters.

In addition to uncertainty arising from incomplete data, many biological processes modify molecular topology through operations that are mathematically equivalent to attaching or removing bands. Enzymes such as topoisomerases and site-specific recombinases alter DNA configuration by cutting, twisting, and rejoining strands, producing controlled topological changes that closely paral-

lel band surgeries in knot theory [10,11]. Such reconnection events can generate or remove crossings, change knot or link type, or create composite structures from simpler precursors, and therefore represent natural biological analogues of band attachments. More broadly, the dynamic organization of chromatin and other polymeric biomolecules involves local strand–passage and loop–formation events that likewise resemble band moves at a coarse topological level [12]. Despite the ubiquity of these band–like transformations in molecular biology, the mathematical theory of *banded knots* has not been systematically developed in this context. This gap suggests an opportunity to integrate band surgery with topological data analysis, enabling new tools for detecting, classifying, and quantifying biologically relevant topological modifications in settings where experimental resolution is limited.

Research Objectives and Proposed Approach

The goal of this project is to develop a new mathematical and computational framework for analyzing knot-like structures in settings where both topological uncertainty and biologically meaningful modifications must be taken into account. Building on the foundations of topological data analysis [13] and the algebraic theory of multiparameter persistence modules [14], we propose to construct a biparameter persistence model in which a geometric “distance” parameter identifies the underlying knotted core of a structure, while an independent “density” parameter reveals additional strands, attachments, or band–like features that may arise from enzymatic activity or incomplete structural reconstruction. This bifiltration will serve as the basis for a theory of *persistent Khovanov homology*, extending both the classical Khovanov homology of links [15] and its behavior under controlled knot operations [16] into a parameterized and data-driven setting. By drawing on the stability principles that underpin persistent homology [17], our objective is to establish a framework in which Khovanov-theoretic information can be tracked across spatial scales and through biologically motivated band surgeries, enabling the detection and classification of knot-like configurations even when full geometric information is unavailable or when the topology itself may change.

To develop this framework, we introduce a distance–density bifiltration that captures two complementary aspects of knot-like structures. The first parameter, *distance*, governs a filtration in which points or segments of an embedded curve are incorporated according to their geometric proximity, allowing a simplified knotted core to emerge across scales. The second parameter, which we refer to as a *density* parameter, governs the addition of band attachments between arcs that lie sufficiently close in the ambient space. At low density levels, the filtration contains only the most reliably supported portions of the structure, while higher density thresholds allow geometrically proximate arcs to be joined by canonical band moves. Such a band move corresponds to a saddle cobordism, and therefore induces a well-defined map (up to sign) on Khovanov homology by the functoriality results of Jacobsson [18]. In this way, the density parameter generates a structured family of knots and banded knots connected by cobordism-induced maps, providing a natural setting in which to examine how Khovanov homology evolves along a bifiltration. This conceptual foundation leads directly to the research aims outlined below.

Specific Aims

The development of persistent Khovanov homology for knots and banded knots requires advances in both the theoretical foundations of bifiltrations and the algebraic structure of Khovanov-type invariants. Guided by the distance–density framework described above, our project will pursue the following specific aims:

1. **Formalize the distance–density bifiltration for knots and banded knots.** We will develop a rigorous mathematical model in which geometric proximity and band attachments are encoded as a two-parameter family of inclusions. This includes identifying conditions under which band additions arise canonically from density thresholds and analyzing how these operations interact with classical knot invariants.
2. **Construct persistent Khovanov homology along the bifiltration.** Using the functoriality of Khovanov homology under saddle cobordisms [18], we will define homomorphisms associated to band moves and assemble these into a multiparameter persistence module. Our goal is to characterize the resulting algebraic structures, study their dependence on filtration parameters, and identify features that persist across scales or through band attachments.
3. **Investigate computational and structural properties of the resulting invariants.** We will implement computational experiments to explore how persistent Khovanov homology behaves for families of knots and banded knots generated by the bifiltration. This includes analyzing small examples, examining how persistent features correspond to known topological operations, and identifying patterns or conjectures that may guide future theoretical development.

These aims establish a coherent pathway toward a persistent version of Khovanov homology adapted to settings in which topological modifications arise naturally or in which partial or uncertain structural information must be incorporated. Together, they lay the groundwork for a broader theory of categorified invariants in multifiltration contexts.

References

- [1] A. Stasiak and J. R. Lúcios, “The why and how of DNA unlinking,” *Nucleic Acids Research*, vol. 37, no. 3, pp. 661–671, 2009.
- [2] A. D. Bates and J. M. Berger, “Keeping intracellular DNA untangled: A new role for topoisomerases,” *BioEssays*, vol. 44, no. 12, p. 2100187, 2022.
- [3] P. F. N. Faísca, “Knotted proteins: A tangled tale of structural biology,” *Journal of the Royal Society Interface*, vol. 12, no. 110, p. 20141182, 2015.
- [4] J. Smrek, D. Michieletto, D. Marenduzzo, and A. Stasiak, “Epigenetic transitions and knotted solitons in stretched chromatin,” *Scientific Reports*, vol. 7, no. 1, pp. 1–11, 2017.
- [5] M. J. Boniecki, G. Lach, W. Dawson, and J. M. Bujnicki, “Entanglements of structure elements revealed in RNA 3D models,” *Nucleic Acids Research*, vol. 49, no. 17, pp. 9625–9642, 2021.
- [6] R. Henderson, “Avoiding the pitfalls of single particle cryo-electron microscopy: Einstein from noise,” *Quarterly Reviews of Biophysics*, vol. 46, no. 2, pp. 135–158, 2013.
- [7] J. Dekker, M. A. Marti-Renom, and L. A. Mirny, “The 3d genome as moderator of chromosomal communication,” *Nature Reviews Genetics*, vol. 14, no. 12, pp. 790–803, 2013.
- [8] F. Benedetti, J. Dorier, Y. Burnier, and A. Stasiak, “Predicting chromosome topological entanglement using polymer models,” *Proceedings of the National Academy of Sciences*, vol. 111, no. 21, pp. E2157–E2164, 2014.

- [9] K. C. Millett, A. Dobay, and A. Stasiak, “Knot localization in polymer chains,” *Macromolecules*, vol. 38, no. 2, pp. 601–606, 2005.
- [10] J. C. Wang, “Dna topoisomerases,” *Nature*, vol. 384, no. 6600, pp. 11–13, 1996.
- [11] N. J. Crisona, R. Weinberg, B. J. Peter, D. W. L. Sumners, and N. R. Cozzarelli, “Processive recombination by phage λ integrase: insights into the mechanism of dna strand exchange,” *Cell*, vol. 99, no. 5, pp. 553–562, 1999.
- [12] A. Stasiak and D. W. L. Sumners, “Mathematics of dna entanglement,” *Science*, vol. 239, no. 4846, pp. 293–299, 1988.
- [13] G. Carlsson, “Topology and data,” *Bulletin of the American Mathematical Society*, vol. 46, no. 2, pp. 255–308, 2009.
- [14] G. Carlsson and A. Zomorodian, “The theory of multidimensional persistence,” *Discrete and Computational Geometry*, vol. 42, no. 1, pp. 71–93, 2009.
- [15] M. Khovanov, “A categorification of the jones polynomial,” *Duke Mathematical Journal*, vol. 101, no. 3, pp. 359–426, 2000.
- [16] P. Turner, “Calculating bar-natan’s characteristic two khovanov homology,” *Journal of Knot Theory and Its Ramifications*, vol. 15, no. 10, pp. 1335–1356, 2006.
- [17] D. Cohen-Steiner, H. Edelsbrunner, and J. Harer, “Stability of persistence diagrams,” *Discrete and Computational Geometry*, vol. 37, no. 1, pp. 103–120, 2007.
- [18] M. Jacobsson, “An invariant of link cobordisms from khovanov homology,” *Algebraic & Geometric Topology*, vol. 4, pp. 1211–1251, 2004.