

# NumbaCS: A fast Python package for coherent structure analysis

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## Summary

NumbaCS (Numba Coherent Structures) is a Python package that efficiently implements a variety of methods for studying material transport in time-dependent fluid flows. It leverages Numba – a high performance Python compiler for generating optimized machine code from Python functions – along with other Numba-compatible packages behind the scenes, producing fast and user-friendly implementations of coherent structure methods. “Coherent structure methods” refer to any method which can be used to infer or extract Lagrangian and objective Eulerian coherent structures. The theory behind these methods have been developed over the last few decades with the aim of extending many of the important invariant objects from time-independent dynamical systems theory to the more general setting where a system may have arbitrary time dependence and may only be known or defined for some finite time. These time-dependent systems are ubiquitous in the context of geophysical and engineering flows where the evolution of the velocity field depends on time and velocity data representing these flows is not available for all time. By extending the ideas from the time-independent setting to the more general time-dependent setting, important transient objects (coherent structures) can be identified which govern how material is transported within a flow. Understanding material transport in flows is of great importance for applications ranging from monitoring the transport of a contaminant in the ocean or atmosphere to informing search and rescue strategies for persons lost at sea.

## Statement of need

As theory and implementations of coherent structures have been developed ([Farazmand & Haller, 2012](#); [Haller, 2011](#); [Haller et al., 2016](#); [Haller & Beron-Vera, 2013](#); [Haller & Poje, 1998](#); [Mathur et al., 2007](#); [Nolan, Serra, et al., 2020](#); [Schindler et al., 2012](#); [Serra & Haller, 2016](#); [Shadden et al., 2005](#)) and the utility of these tools has been demonstrated over the last two decades ([Curbelo & Rypina, 2023](#); [Du Toit & Marsden, 2010](#); [Günther et al., 2021](#); [Liu et al., 2018](#); [Nolan, Foroutan, et al., 2020](#); [Peacock & Haller, 2013](#); [Pretorius et al., 2023](#); [Rutherford et al., 2012](#); [Serra et al., 2017](#)), there has been a steadily growing interest in using these methods for real-world applications. Early on, software implementations were largely contained to in-house packages developed by applied mathematicians and engineers advancing the theory. Over the years, there have been a number of software packages developed in an attempt to provide implementations of some of these methods for practitioners outside of the field. Some provide a friendly interface for users (Dynlab – [Nolan \(2024\)](#); LCS MATLAB Kit – [Dabiri \(2009\)](#)), others aim to provide efficient implementations of specific methods (sometimes in specific circumstances) (Lagrangian – [Briol & d’Ovidio \(2011\)](#); Newman – [Du Toit \(2010\)](#); Aquila-LCS – [Lagares & Araya \(2023\)](#)), and a few implement a variety of methods (Tbarrier – [Bartos et al. \(2022\)](#); LCS Tool – [Onu et al. \(2015\)](#)). NumbaCS intends to unite these aims by providing efficient and user-friendly implementations of a variety of coherent structure

methods. By doing this, the hope is to provide a powerful tool for experienced practitioners and a low barrier of entry for newcomers. In addition, as new methods/implementations arise, the framework laid out in NumbaCS provides a straightforward environment for contributions and maintenance. Also of note is another package called `CoherentStructures.jl` ([Junge et al., 2020](#)), which is fast, user-friendly, and implements a variety of methods. This package has some overlap with NumbaCS but they both implement methods which the other does not. `CoherentStructures.jl` is a powerful tool that should be considered by users who perhaps prefer Julia to Python or are interested in computing some of the methods not implemented in NumbaCS. For a more detailed breakdown of how all of the mentioned packages compare with NumbaCS, see the [documentation](#).

## Functionality

NumbaCS implements the following features for both analytical and numerical flows:

- Standard flow map computation
- Flow map composition method ([Brunton & Rowley, 2010](#))
- Finite time Lyapunov exponent (FTLE) ([Shadden et al., 2005](#))
- instantaneous Lyapunov exponent (iLE) ([Nolan, Serra, et al., 2020](#))
- Lagrangian averaged vorticity deviation (LAVD) ([Haller et al., 2016](#))
- Instantaneous vorticity deviation (IVD) ([Haller et al., 2016](#))
- FTLE ridge extraction ([Schindler et al., 2012](#); [Steger, 1998](#))
- Variational hyperbolic LCS ([Farazmand & Haller, 2012](#); [Haller, 2011](#))
- Variational hyperbolic OECS ([Serra & Haller, 2016](#))
- LAVD-based elliptic LCS ([Haller et al., 2016](#))
- IVD-based elliptic OECS ([Haller et al., 2016](#))

For flows defined by numerical velocity data:

- Simple creation of JIT compiled linear and cubic interpolants

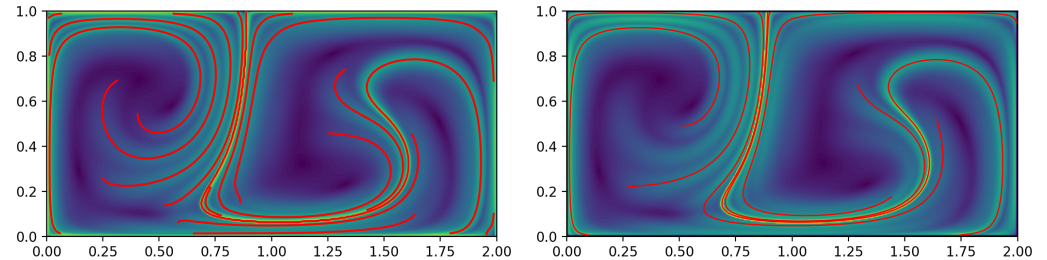
All of these implementations are relatively straightforward to use and quite efficient. This is due to three key dependencies NumbaCS utilizes to speed up computations. The first is Numba ([Lam et al., 2015](#)), a JIT compiler for Python which can drastically speed up numerical operations and provides a simple framework for parallelizing tasks. Next, `numba_ode` ([Wogan, 2021](#)) is a Python wrapper to ODE solvers in both C++ (LSODA) and FORTRAN (DOP853) that bypasses the Python interpreter and can be used within Numba functions (common Python ODE solvers, such as those provided by the `SciPy` package, cannot be executed within Numba functions). This package is crucial to the efficiency of NumbaCS as particle integration is often the most costly part of finite-time coherent structure methods. Finally, the interpolation package ([Winant et al., 2017](#)) provides optimized interpolation in Python and is utilized in NumbaCS to create JIT compiled interpolant functions, producing efficient implementations of methods even for flows defined by numerical data. By taking advantage of these packages behind the scenes, NumbaCS is able to maintain the simplicity and readability of an interpreted language while achieving runtimes closer to that of a compiled language.

## Examples

A [User Guide](#) is provided which details the workflow in NumbaCS and a number of examples demonstrating the functionality are covered in the [Example Gallery](#). Here we show the output of a few examples, provide the runtime of each, and breakdown the runtime based on the parts of each method. “Flowmap” refers to the particle integration step, “C eig” and “S eig” refer to the eigenvalue/vector step for Lagrangian and Eulerian methods respectively (this time will be roughly equal to the FTLE and iLE times), and the last is the extraction time for a given method. For examples that require particle integration, the default solver (DOP853) was used

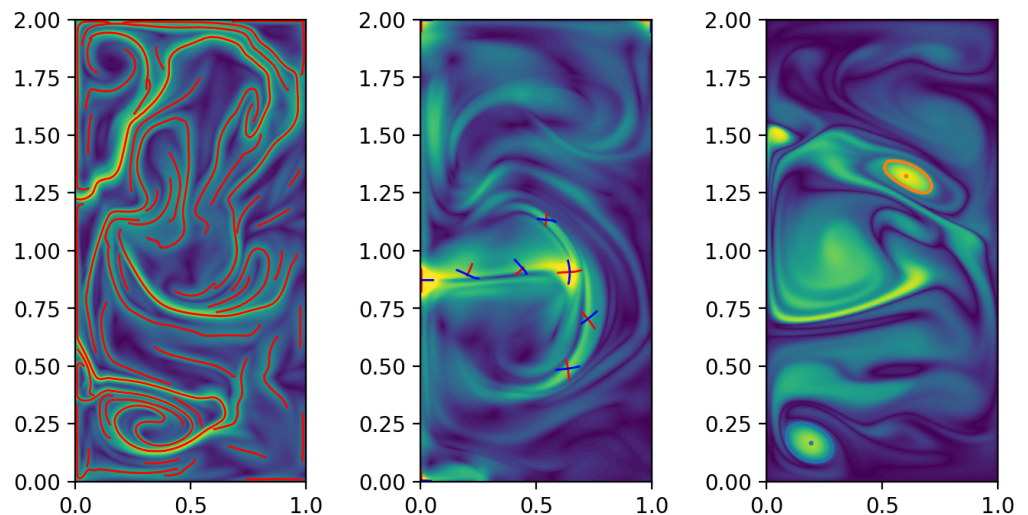
with the default error tolerances (relative tolerance =  $1e-6$ , absolute tolerance =  $1e-8$ ). All runs were performed on an Intel<sup>(R)</sup> Core<sup>TM</sup> i7-3770K CPU @ 3.50GHz (which has 4 cores and 8 total threads). Warm-up time<sup>1</sup> is not included in the timings.

### Analytical Flow (Double Gyre)



Left: **DG FTLE ridges** at  $t_0 = 0$ , integration time  $T = -10$ . Total runtime per iterate:  $\sim 0.424s$  (flowmap:  $\sim 0.390s$ ; C eig:  $\sim 0.025s$ ; FTLE ridge extraction:  $\sim 0.009s$ ). Right: **DG hyperbolic LCS** at  $t_0 = 0$ , integration time  $T = -10$ . Total runtime per iterate:  $\sim 5.219s$  (flowmap (aux grid):  $\sim 1.83s$ ; C eig (aux grid):  $\sim 0.039s$ ; hyperbolic LCS extraction:  $\sim 3.350s$ ). Both are computed over a  $401 \times 201$  grid.

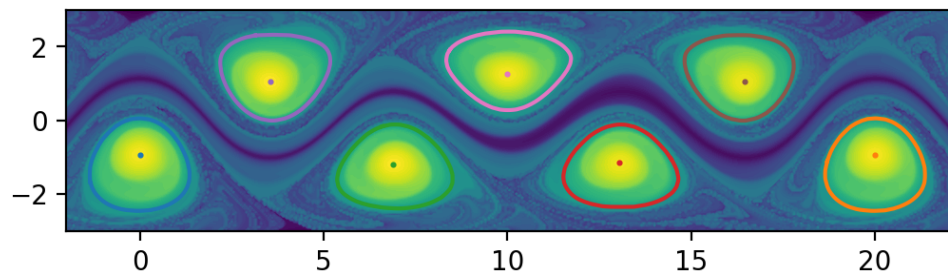
### Numerical Flow (QGE)



Left: **QGE FTLE ridges** at  $t_0 = 0$ , integration time  $T = 0.1$ . Total runtime per iterate:  $\sim 2.461s$  (flowmap:  $\sim 2.400s$ ; C eig:  $\sim 0.038s$ ; FTLE ridge extraction:  $\sim 0.023s$ ). Middle: **QGE hyperbolic OECS** at  $t_0 = 0.15$ . Total runtime per iterate:  $\sim 2.238s$  (S eig:  $\sim 0.038s$ ; hyperbolic OECS extraction:  $\sim 2.200s$ ). Right: **QGE elliptic OECS** at  $t_0 = 0.5$ . Total runtime per iterate:  $\sim 0.0452s$  (IVD:  $\sim 0.0002s$ ; elliptic OECS extraction:  $\sim 0.045s$ ). All are computed over a  $257 \times 513$  grid.

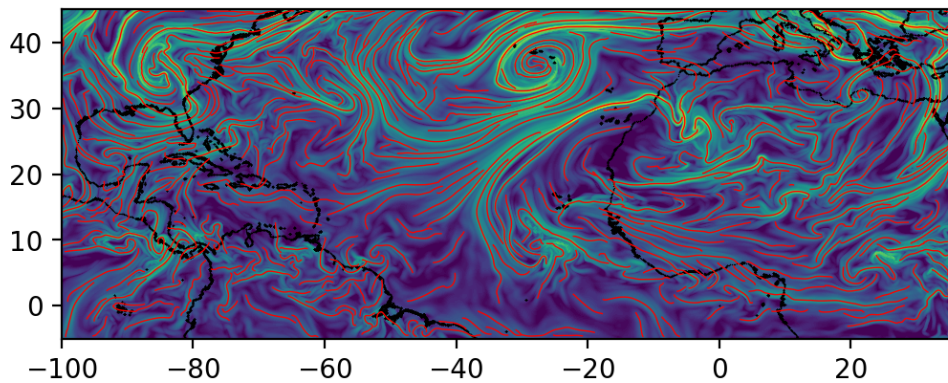
<sup>1</sup>Since many functions in NumbaCS are JIT compiled, these functions are optimized and compiled into machine code on the first function call. This initial delay is often referred to as “warm-up time”. After the first call, subsequent function calls are much faster.

### Analytical Flow (Bickley jet)



**Bickley jet elliptic LCS** at  $t_0 = 0$ , integration time  $T = 40$  days. Total runtime per iterate:  $\sim 5.065$ s (flowmap:  $\sim 4.490$ s; LAVD:  $\sim 0.565$ s; elliptic LCS extraction:  $\sim 0.010$ s). Computed over  $482 \times 121$  grid.

### Numerical Flow (MERRA-2)



**MERRA-2 FTLE ridges** at  $t_0 = 06/16/2020-00:00$ , integration time  $T = -72$ hrs. Total runtime per iterate:  $\sim 7.835$ s (flowmap:  $\sim 7.480$ s; C eig:  $\sim 0.085$ s; FTLE ridge extraction:  $\sim 0.27$ s). Computed over  $902 \times 335$  grid.

## Datasets

Two datasets are provided with NumbaCS to test the functionality for flows defined by numerical velocity data. One is a numerical simulation of the quasi-geostrophic equations (QGE). We thank the authors of Mou et al. (2021) for providing us with this dataset, which was used extensively during development, and allowing a piece of the dataset to be included in the package. The full dataset was over the time span  $[10, 81]$  with  $dt = 0.01$ . We provide the velocity fields over the much shorter time span of  $[10, 11]$  with the same  $dt$ . For details on parameters used in the simulation, refer to the cited paper. The other dataset is a MERRA-2 vertically averaged reanalysis dataset (Gelaro et al., 2017; GMAO, 2015), which was used as part of a paper (Jarvis et al., 2024) coauthored by the authors of this paper. Wind velocity fields were vertically averaged over pressure surfaces ranging from 500 hPa to 800 hPa. The corresponding latitude, longitude, and date arrays are also provided. All data can be downloaded from the [data folder](#) on the GitHub page.

## Usage in ongoing research

As of the writing of this paper, NumbaCS has not been public for long but has been utilized in one publication (Jarvis et al., 2024) where it was the computational tool for all coherent



structure methods. In addition, it is currently being used in an ongoing project focused on airborne invasive species traveling from Australia to New Zealand titled “[Protecting Aotearoa from wind-dispersed pests](#)”. This is a five year (October 2023 - October 2028) Scion-led and Ministry of Business, Innovation and Employment (MBIE)-supported program.

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