




SODA: Software to Support the Curation and Sharing of FAIR Autonomic Nervous System Data

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Software

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Summary

SODA (Software to Organize Data Automatically) is an open source and free cross-platform desktop software that assists researchers in preparing and sharing their autonomic nervous system (ANS) related data according to the guidelines developed by the National Institute of Health (NIH)'s [Stimulating Peripheral Activity to Relieve Conditions \(SPARC\) Program](#). By combining intuitive user interfaces with automation, SODA streamlines the process of implementing the SPARC guidelines which can otherwise be challenging and/or time consuming for researchers.

Statement of need

SPARC was established in 2014 to accelerate the development of therapeutic devices that modulate electrical activity in the ANS to improve organ function. Research on bioelectronic medicine had shown tremendous potential to address diverse conditions such as hypertension, heart failure, and gastrointestinal disorders but required further investigation and development for clinical translation, which SPARC aimed to support. In addition, the SPARC Program also supported the development of guidelines for making data resulting from such research optimally reusable in line with the FAIR (Findable, Accessible, Interoperable, Reusable) Principles ([Wilkinson et al., 2016](#)). This includes microscopy imaging, electronic measurement data, and more. The goal was to provide a standard way for researchers to organize and share their ANS-related data to facilitate secondary data reuse, enable joint analysis with different datasets, and accelerate discoveries ([Quey et al., 2021](#); [Soundarajan et al., 2022](#)). Accordingly, the [SPARC data curation and sharing guidelines were developed](#). The guidelines prescribe the data to be organized according to the standard SPARC Data Structure (SDS) ([Bandrowski et al., 2021](#)) and shared publicly on [sparc.science](#), the data portal of the SPARC program ([Osanlouy et al., 2021](#)). The SPARC data curation and sharing guidelines have been imposed on all SPARC-funded researchers since 2017. Since 2022, [sparc.science](#) has become an open repository so anyone with ANS-related data can follow the SPARC guidelines to have their dataset published on the SPARC portal. The guidelines are very exhaustive to maximize FAIRness of data but as a result, they are challenging and time-consuming for researchers to implement. While being part of a SPARC-funded research project in 2017, we experienced these challenges firsthand. The process to implement the SPARC guidelines involves several data manipulations to comply with the SDS, which includes organizing data according to a strict folder structure, following file and folder naming conventions, preparing several metadata files, using standard formats for certain data types, and so on. Then, the resulting dataset needs to be uploaded on [Pennsieve](#), the data management platform of the SPARC Program. There, the dataset is reviewed by a team of human curators and a back-and-forth conversation could ensue to address any SDS-compliance issues before the dataset is published on [sparc.science](#).

To simplify this process for the data resulting from our SPARC-funded project, we developed Python scripts to automate some of the steps. We presented this automation approach during a SPARC-organized Hackathon in 2018, where it received wide appreciation from other SPARC-funded researchers who were experiencing similar challenges. The project won the Public's Choice Award at the Hackathon and subsequently received funding from the SPARC Program starting in 2019 for further development of this idea into a desktop software application that anyone can use without coding knowledge. This gave birth to SODA, the software presented in this paper (Patel et al., 2020).

Software overview

SODA is a cross-platform desktop software application developed using [Electron](#), an open-source framework for creating desktop applications using web technologies. [Flask](#) is used in the backend of the software to integrate with existing tools that help with complying with the SPARC guidelines, which are mostly developed in Python. Further information about the technical development of SODA can be found in its GitHub repository (Patel, Soundarajan, Marroquin, et al., 2023). SODA is designed to guide researchers step-by-step through all the requirements for preparing and sharing their data according to the SPARC guidelines. The software combines intuitive user interfaces with automation to streamline the process. A screenshot of the home page of SODA is provided in [Figure 1](#). The development was started in 2019 and is still ongoing as support for more elements of the SPARC guidelines along with additional automation are progressively being implemented. SODA integrates with other SPARC resources such as the SDS validator (Gillespie, 2023), the [Pennsieve API](#), and more integrations are planned such that SODA becomes a one-stop tool for anyone wanting to make their ANS-related data FAIR through the SDS and the SPARC data portal. A full overview of what SODA supports is available in its [user documentation](#).

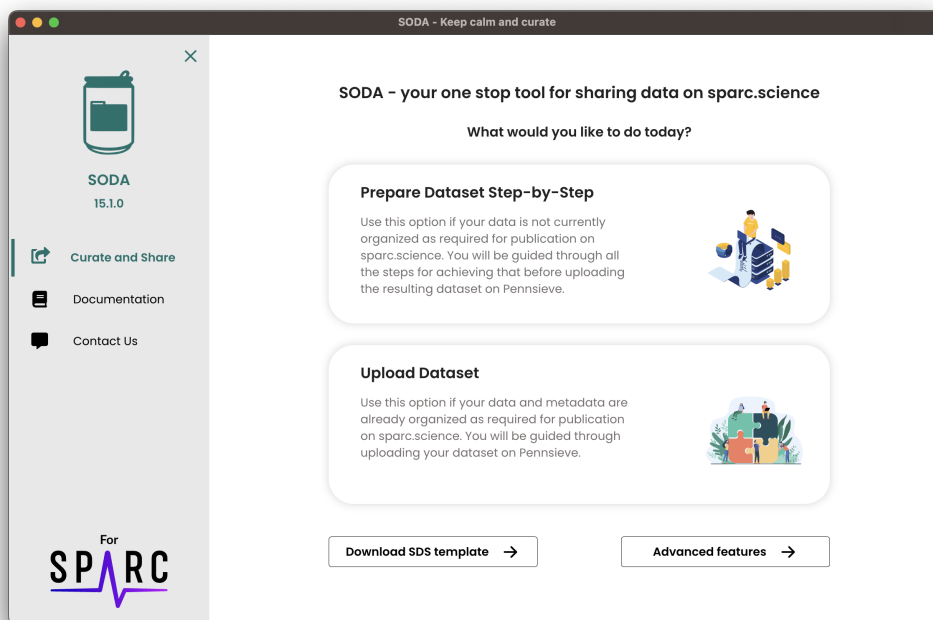


Figure 1: Screenshot of the user interface of the home page of SODA.

Performance, usage, and impact

Beta testing was conducted in 2020 after about one year of developing SODA to evaluate its performance. The beta testers consisted of 10 SPARC-funded researchers from 10 different research groups who had never used SODA before but some of them had prepared and submitted data according to the SPARC guidelines without SODA. They were provided a sample dataset and asked to prepare and submit it as per the SPARC guidelines without SODA (Task A) and with only SODA (Task B). For both tasks, they were asked to complete only the steps of the SPARC guidelines supported by SODA at that time (implementing the SDS folder structure, preparing certain metadata files, uploading to Pennsieve, etc.). Half of them (randomly selected) were asked to complete Task A first then Task B while the other half were asked to complete the tasks the other way around. They were requested to report back the time required to complete each task and score out of 5 (1: very difficult, 5: very easy) the ease of understanding and implementing the SPARC guidelines for each task. Overall, we found that SODA reduced the time required to prepare and share a dataset according to the SPARC guidelines by 70% and made it relatively easier to understand and implement the requirements (Figure 2). After evaluating the shared datasets on Pennsieve, we found that there was an aggregate of 23 compliance errors in datasets submitted without SODA while only one error was found amongst datasets submitted with SODA. When subsequently asked if they would consistently prepare and share their data (SPARC or else) if not mandatory, a majority (8/10 yes, 2/10 maybe) responded that they would if a software application like SODA was available while a majority would not without it (6/10 no, 4/10 maybe). After improving the user flow, adding more automation, and additional integration with other SPARC resources since that beta testing, we believe that SODA is reducing researchers' time, effort, and error in the process even more now.

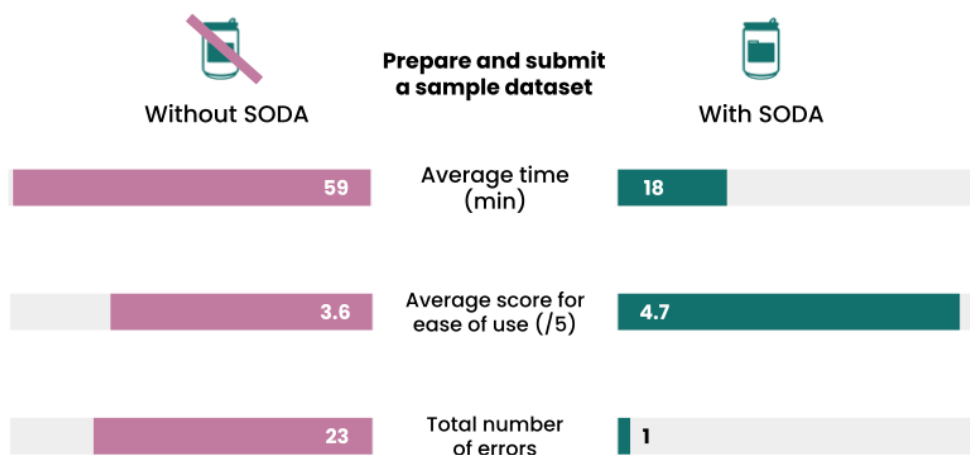


Figure 2: Results from testing of SODA by 10 beta testers during 2020.

Given these advantages offered by SODA, it has been widely used (over 2,000 downloads) by SPARC researchers since 2020 and by researchers collecting ANS-related data outside of SPARC since 2022, when sparc.science became an open repository. Since the beginning of 2021, SODA has helped researchers all over the world process over 27TB of data corresponding to over 400k individual data files (as of May 2024). The development of SODA is still on-going to further simplify the process of formatting datasets according to the SDS and publishing them on the SPARC data portal. Additionally, there are several software engineering improvements that need to be made, such as adding an authentication guard for the Flask server and removing duplicate constants. They are anticipated to be completed by the end of the next SODA grant cycle in August 2025.

There is a major push to make data FAIR in all fields of research, including biomedical research with the NIH's leadership. As a result, many standards, guidelines, and platforms to archive data are developed to achieve that. However, the burden of understanding, learning, and using these resources for making data FAIR is mostly left to the researchers. To our knowledge, SODA is the first researcher-oriented tool that guides step-by-step through all the requirements for making data FAIR from organizing files, creating metadata, validating, and uploading to a data repository. It has since inspired several other tools we are developing such as FAIRshare (Soundarajan & Patel, 2023) through support from the National Institute of Allergy and Infectious Diseases (NIAID) and fairhub.io (Soundarajan et al., 2023) through support from the NIH Bridge2AI Program. The codebase of SODA was also forked by another team that is developing a tool called NWB GUIDE to simplify the process of preparing and sharing data from the NIH Brain Initiative Program. SODA has itself been made FAIR in line with the FAIR-BioRS guidelines (Patel, Soundarajan, Ménager, et al., 2023) to promote and facilitate such reuse of its source code outside of the developing team.

Acknowledgements

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