

Scientific Paper

Benjamin Vandersmissen¹, Lars Van Roy²,
Evelien Daems³, and Frank Jan Fekkes⁴

¹ `benjamin.vandersmissen@student.uantwerpen.be`

² `lars.vanroy@student.uantwerpen.be`

³ `evelien.daems@student.uantwerpen.be`

⁴ `franciscus.fekkes@student.uantwerpen.be`

Abstract. This paper will give an in depth view of the performed adaptations to the Stride project, as well as an overview of the findings that were obtained by comparing these additions to the original stride project.

1 Introduction

The stride project is designed to simulate and evaluate the lifetime of various infectious diseases. In order to properly analyze which factors have an effect on various diseases, the stride project allows multiple factors to be varied so that we can get an idea of which factors are the main influence for the evolution of diseases. This expansion to the original stride project will include 2 major corrections, to make the simulation more realistic, being an improved workplace size distribution and an improved household composition distribution, a major addition to the pools in which diseases can spread, being daycare and preschool pools, a minor addition to make the input more variable by adding two addition input types being HDF5 and JSON and finally a visualizer that will allow users to get a graphical overview of where the diseases are most active, along with a list of parameters to give further information of the evolution of the disease for a given location.

Other than these additions, there will also be a section about the simulation over entire Belgium, compared to isolated within Flanders. Other than just the simulated area, there will also be a comparison between the data derived from the Belgian population and the data derived from the Flemish population.

2 Daycare & Preschool

3 Data Formats

3.1 JSON

3.2 HDF5

HDF5 is a scientific dataformat specifically designed to store and organise large amounts of data. In this section, we compare the sizes of the fileformats already

available in stride (protobuf and JSON) with the size of an HDF5 file and we also cover the write speed. Both these metrics are based on the configuration file *run_generate_default.xml* with only the output file changed.

As shown in table 1, HDF5 is memory-wise between protobuf and JSON. This is because protobuf is a heavily compressed format and HDF5 is not, but HDF5 isn't a plain text format like JSON either. In terms of write speed, HDF5 is a lot slower than Protobuf and JSON. A simple explanation for this is the fact that HDF5 does not support streams, while JSON and Protobuf do. This means that HDF5 has to write everything with it's own procedures which isn't as efficient. HDF5 does not support streams, because it was originally a C library, where streams did not exist and the C++ implementation is just a wrapper around the original C library.

FileFormat	Size (Kb)	Average write Time (s)
Protobuf	13.713	1.74
JSON	317.050	5.25
HDF5	125.066	30.45

Table 1. Performance of different data formats based on 10 runs of *run_generate_default.xml*

4 Data Visualization

5 Workplace Size Distribution

5.1 Introduction

The original stride project made use of a set workplace size distribution, being an average size of 20, that would be randomly populated. So it might be that there would be smaller and larger workplaces, but the average would be around 20, which is not accurate. The majority of the workplaces are smaller than 20 and there are workplaces that are way larger than 20. It would therefore be more accurate to add an input file that would give a distribution of which workplace size occurs with which chance.

5.2 Impact

To proof whether or not this addition had a major impact, and to show the size of its impact, we ran a couple of simulations with different values for *r0*. The resulting graphs are displayed in the annex. Even though the difference isn't huge, we can generally conclude that it most definitely made the spreading go faster. At the end of the simulations, the number of infected people is generally about 4% lower compared to using the newer workplace implementation. We can therefore conclude that the original workplace implementation was indeed incorrect, and gave slightly flawed results.

6 Household Composition Distribution

6.1 Introduction

One more flaw of the original stride project, is that it had no regard for the differences in household size within smaller regions. It used one big input file, that would give the general household configuration over the entire simulation area (in this case Flanders) but there are significant differences between households over the different provinces. We therefore wanted to be able to specify household information for each province.

Other than provinces, there are also significant differences between regular cities and major cities. We therefore added one extra household configuration for major cities, along with a file specifying which cities are considered to be major.

6.2 impact

To proof the impact of this addition, we did the same simulations we did for the workplace distribution addition. These plots, found in the annex, are different to the plots from workplaces in that they are not consistent in the effect that occurred. We can see significant effects in all three plots, but for r_0 equal to 8 we see that the new implementation gives a far lower number of diseased people, where the other two show that the new implementation gives a slightly higher number of diseased people, compared to the original stride, spread over the two, the average is still a drop of 2%. We can conclude that this change was significant and that there was indeed a flaw in the original stride implementation or, in other words, the differences between regions and cities are significant to the simulation.

7 Belgian Simulations

Appendices

A Workplace Size Distribution

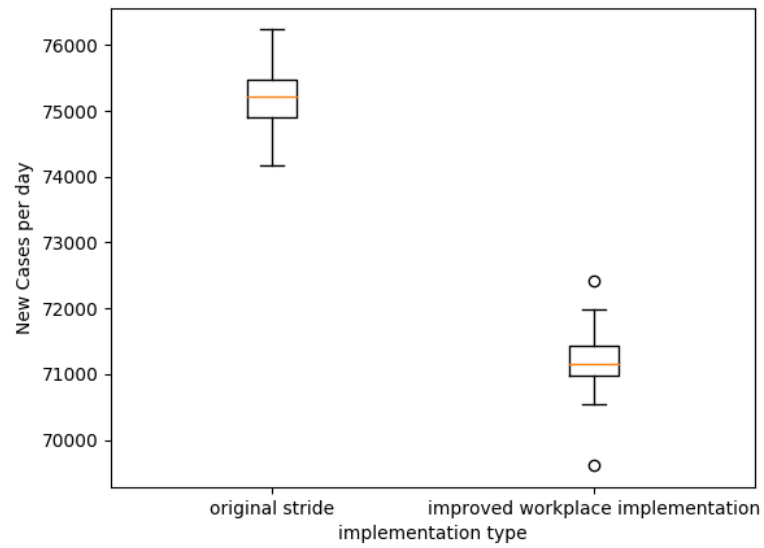


Fig. 1. The boxplot for the different workplace implementations, using r_0 equal to 8

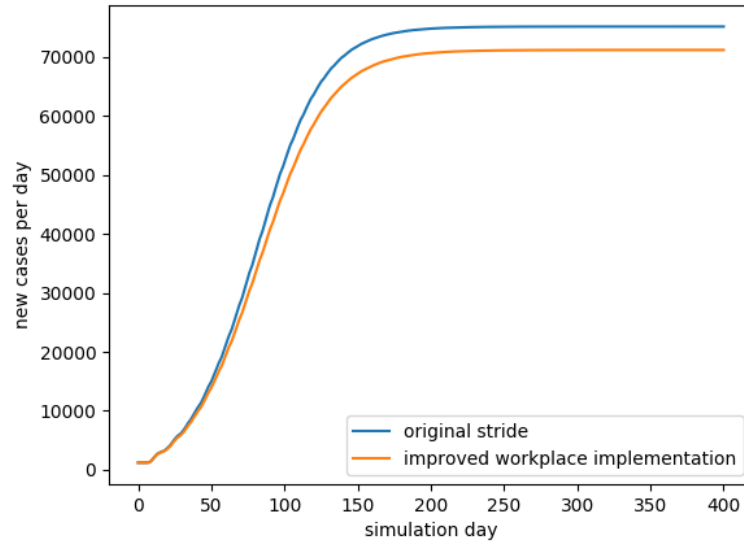


Fig. 2. The cumulative evolution of the diseased people, using the different workplace implementations and r_0 equal to 8

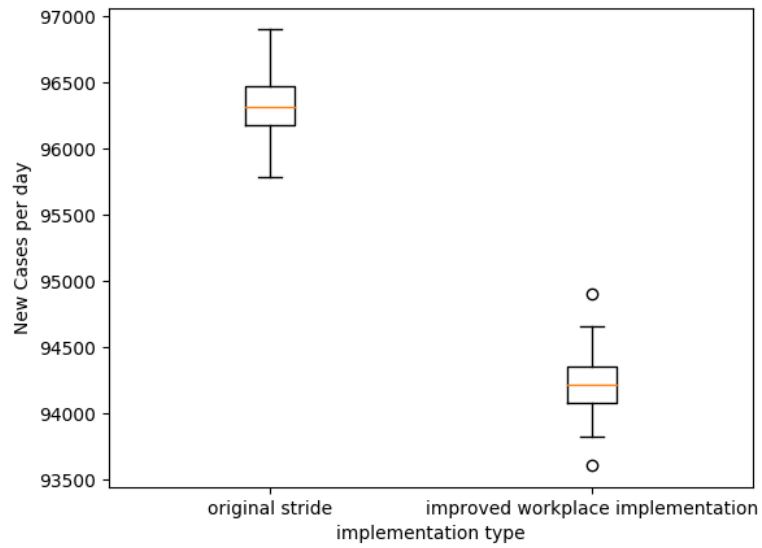


Fig. 3. The boxplot for the different workplace implementations, using r_0 equal to 11

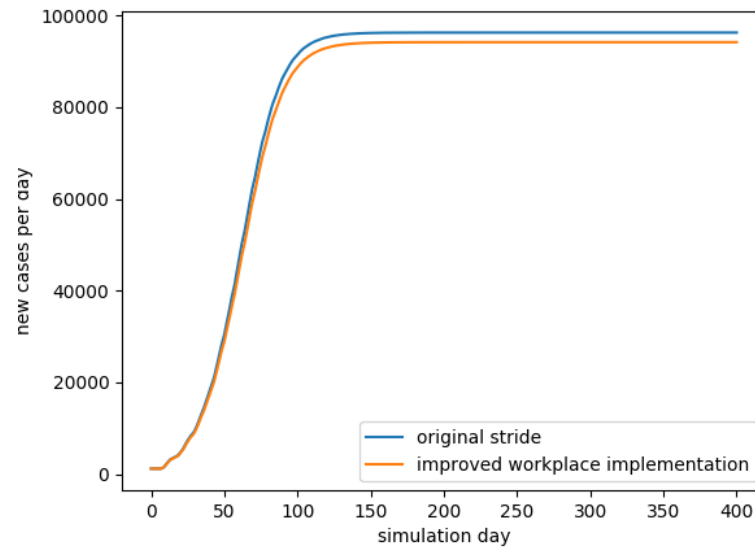


Fig. 4. The cumulative evolution of the diseased people, using the different workplace implementations and r_0 equal to 11

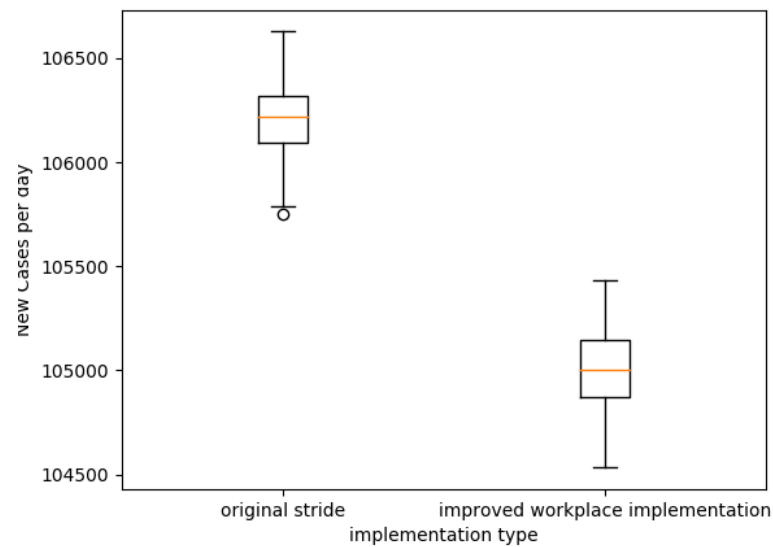


Fig. 5. The boxplot for the different workplace implementations, using r_0 equal to 14

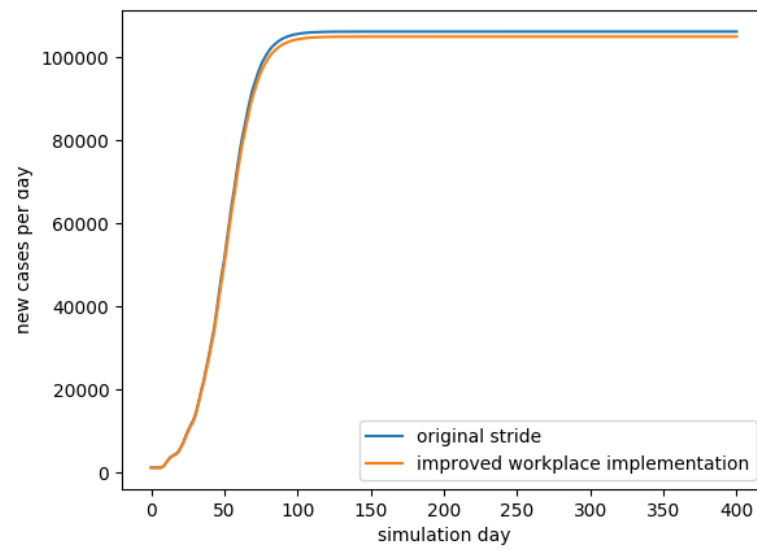


Fig. 6. The cumulative evolution of the diseased people, using the different workplace implementations and r_0 equal to 14

B Household Composition Distribution

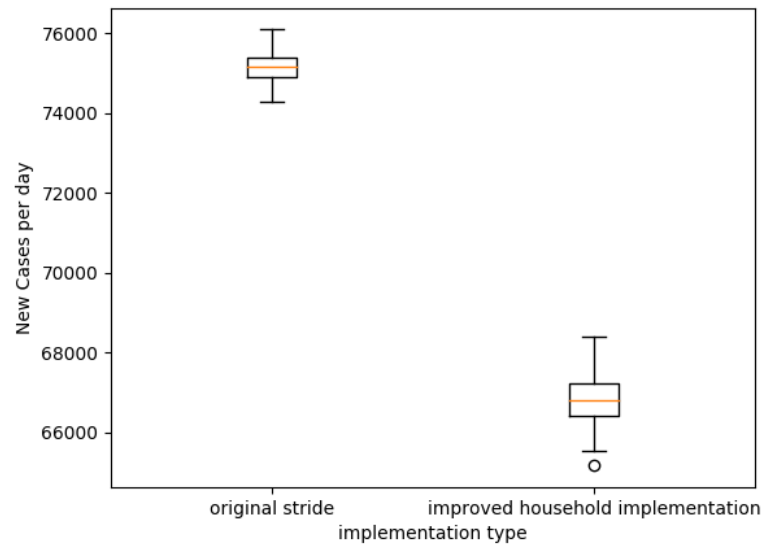


Fig. 7. The boxplot for the different household implementations, using r_0 equal to 8

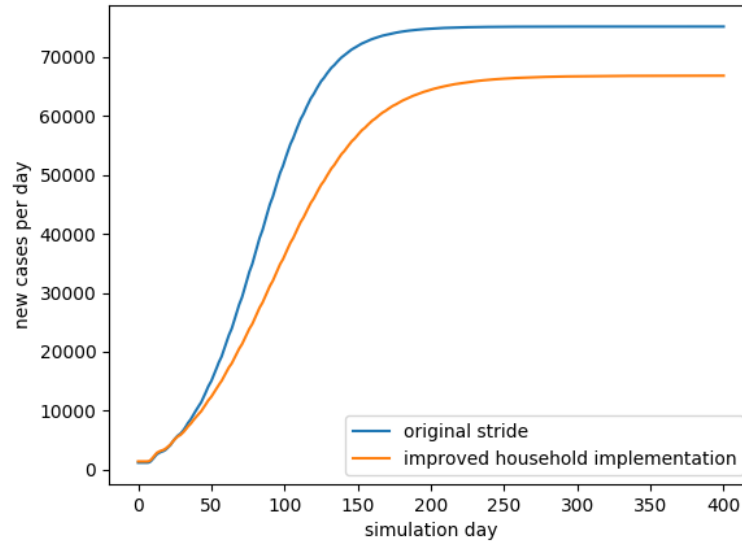


Fig. 8. The cumulative evolution of the diseased people, using the different household implementations and r_0 equal to 8

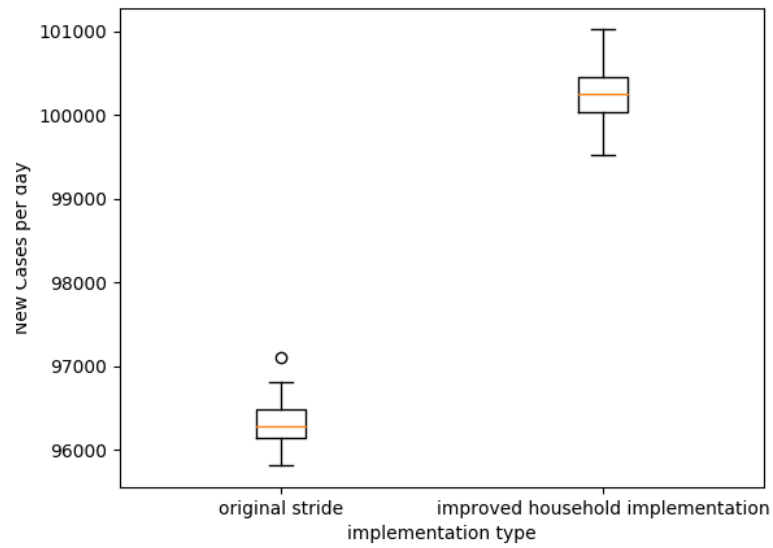


Fig. 9. The boxplot for the different household implementations, using r_0 equal to 11

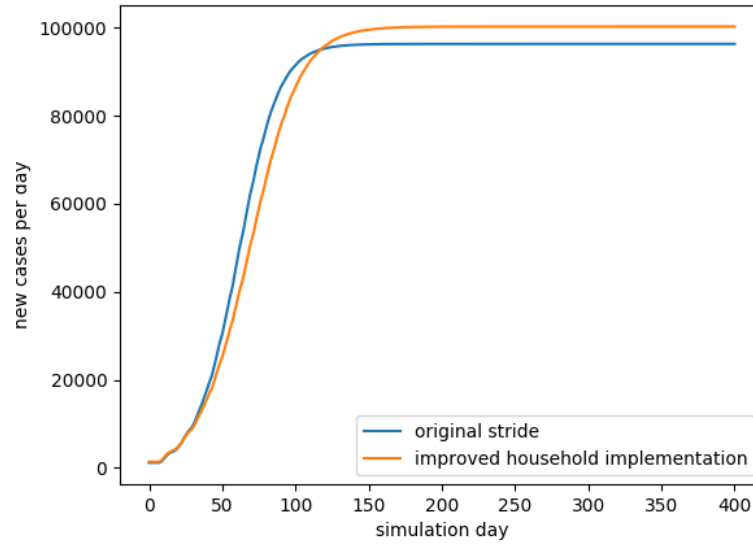


Fig. 10. The cumulative evolution of the diseased people, using the different household implementations and r_0 equal to 11

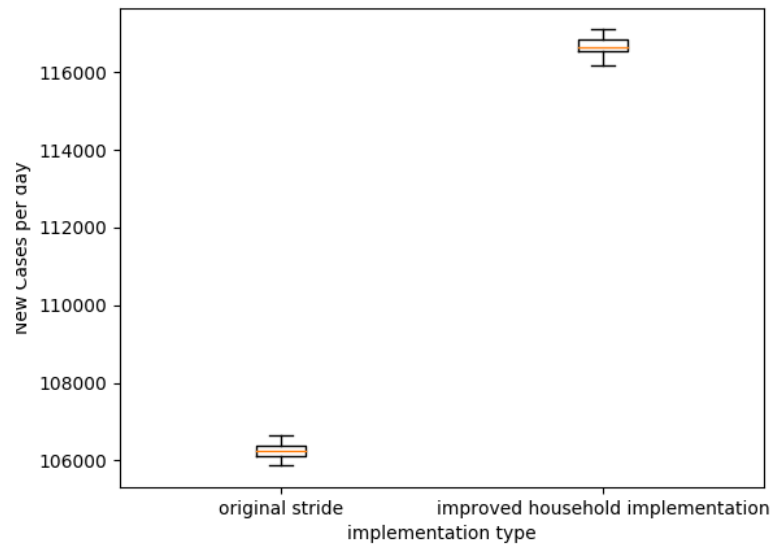


Fig. 11. The boxplot for the different household implementations, using r_0 equal to 14

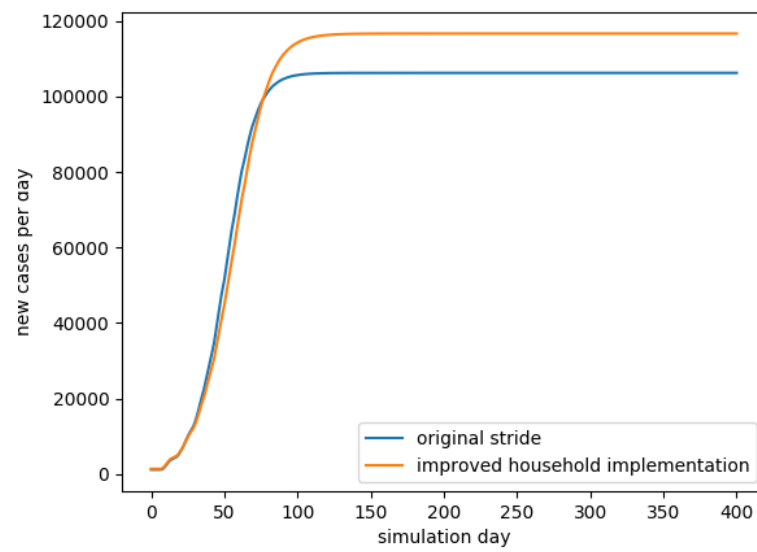


Fig. 12. The cumulative evolution of the diseased people, using the different household implementations and r_0 equal to 14