### TECE progress meeting

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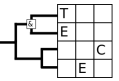
2017-09-13

http://github.com/richelbilderbeek/Science









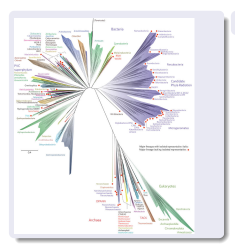
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#### Tree of life<sup>2</sup>



#### Uses RAxML<sup>1</sup>

<sup>&</sup>lt;sup>1</sup>Stamatakis, Alexandros. 2014. Bioinformatics 30.9 (2014).

<sup>&</sup>lt;sup>2</sup>Hug, Laura A., et al. 2016. Nature Microbiology 1 (2016).

# New giraffe species ... 34

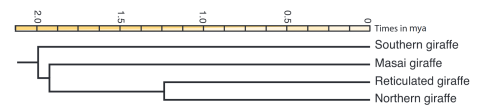


<sup>&</sup>lt;sup>3</sup>Fennessy, Julian, et al. 2016. Current Biology 26.18. <sup>4</sup>Picture by Julian Fennessy



RJC Bilderbeek (University of Groningen)

## ... for quite some time<sup>5</sup>



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## Research question

What is the error we make today ...

- in inferring phylogenies
- caused by ignoring that speciation takes time to complete?

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## From DNA to phylogeny

Using BEAST2<sup>6</sup>Using constant-rate birth-death model<sup>7</sup>

<sup>&</sup>lt;sup>666</sup>Bouckaert, Remco, et al. 2014. PLoS computational biology 10.4.

<sup>666</sup> Nee, Sean, et al. 1994. Philos Trans R Soc Lond B Biol Sci 344.1309. € > € ✓ Q Q

## Simulating DNA

From PBD phylogeny<sup>8</sup> to DNA alignment<sup>9</sup>.

 $<sup>^{666}\</sup>mbox{Etienne},$  Rampal S., and James Rosindell. 2012. Systematic Biology 61.2.

## Measuring error

Using nLTT statistic<sup>10</sup>