

# Identification of Invasive Alien Species using DNA barcodes

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## General introduction to this factsheet

The Barcoding Facility for Organisms and Tissues of Policy Concern (BopCo) aims at developing an expertise forum to facilitate the identification of biological samples of policy concern in Belgium and Europe. The project represents part of the Belgian federal contribution to the European Research Infrastructure Consortium LifeWatch.

Non-native species which are being introduced into Europe, whether by accident or deliberately, can be of policy concern since some of them can reproduce and disperse rapidly in a new territory, establish viable populations and even outcompete native species. As a consequence of their presence, natural and managed ecosystems can be disrupted, crops and livestock affected, and vector-borne diseases or parasites might be introduced, impacting human health and socio-economic activities. Non-native species causing such adverse effects are called Invasive Alien Species (IAS). In order to protect native biodiversity and ecosystems, and to mitigate the potential impact on human health and socio-economic activities, the issue of IAS is tackled in Europe by EU Regulation 1143/2014 of the European Parliament and Council. The IAS Regulation provides for a set of measures to be taken across all member states. The list of Invasive Alien Species of Union Concern is regularly updated. In order to implement the proposed actions, however, methods for accurate species identification are required when suspicious biological material is encountered.

Because morphology-based species identifications are not always possible (e.g. cryptic species, trace material, early life-stages), the purpose of the present work is to investigate and evaluate the usefulness of DNA sequence data to identify each of the IAS included in the EU Regulation. The results are presented as factsheets (one per IAS) compiled using publicly available DNA sequence data and information aggregated from various sources. Each factsheet consists of two major parts; (i) a short introduction to the specific IAS compiling information on its taxonomy and current occurrence/distribution in Europe; (ii) an investigation with respect to the usefulness of publicly available DNA sequences to identify this IAS to the taxonomic level stated in the EU list using DNA barcoding. For further information about the reasoning behind the applied approach and details on the materials and methods utilised, please see below and Smitz *et al.* [1].

More info about BopCo on <u>http://bopco.myspecies.info/</u> or contact us via <u>bopco@naturalsciences.be</u>. More info on the EU Regulation on <u>http://ec.europa.eu/environment/nature/invasivealien/index\_en.htm</u>.

## Salvinia molesta

D.S. Mitch., 1972

Common names: English: African payal, African pyle, Australian azolla, Kariba weed, giant azolla, water fern, giant salvinia, aquarium watermoss, salvinia moss French: salvinia géante, fougère d'eau German: Bueschelfarn, Lästiger Schwimmfarn Dutch: grote vlotvaren



Last update: January 2022

## General information on Salvinia molesta

## Classification

Kingdom	Phylum	Class	Order	Family	Genus					
Plantae	Pteridophyta	Polypodiopsida	Salviniales	Salviniaceae	Salvinia					

## Species in the same genus: N = 10-15 [2-4]

Note: *Salvinia molesta* and three other species are part of a complex, the other species are *S. auriculata, S. biloba* and *S. herzogii*. Morphological uniformity and hybrid origins of these and other species create confusion about the number of extant species. Hybridisation between species of the genus occurs; *S. molesta* itself has intermediate characteristics that would be expected between *S. biloba* and *S. herzogii*.

## Infra-species level: N = 0

Note: To our knowledge, no subspecies have been described.



**Native range:** [5] South-eastern Brazil.

## Invasive range: [6, 7]

Europe (geographical):

Austria, Belgium, Denmark, France, Germany, Italy, Netherlands, Portugal, Romania, Spain, Switzerland.

For more detailed locality information and the most recent distribution updates, please visit:

https://easin.jrc.ec.europa.eu/spexplorer/ https://www.gbif.org/species/5274863 https://gd.eppo.int/taxon/SAVMO/distribution

## Outside Europe (geographical):

Globally widespread.

#### Morphology, biology, invasion, negative effects and remedies

For more information on *Salvinia molesta* please see the references and online information listed at the end of this document.

## Species identification based on DNA barcodes

## Introduction

DNA barcoding is a species identification method that uses a short genetic sequence (DNA barcode) to compare an unknown sample to a database of reference sequences with known species affiliations. The underlying rationale is that the divergence of nucleotide sequences among different species is larger than the nucleotide divergence between sequences within a species. DNA barcoding can facilitate the identification of IAS samples, especially when morphological characteristics are absent or useless. To assure correct species identifications, however, reference libraries need to include a sufficiently large number of sequences of (i) the IAS under investigation, in order to assess the intraspecific genetic divergence; (ii) the closely related species, in order to evaluate the interspecific genetic divergence; (iii) the different geographical areas covering the distribution range (native and invasive) of the IAS in order to detect potential population structure or local hybrids.

Against this background, BopCo evaluated the inclusion of the IAS and their close relatives in both publicly available reference libraries BOLD (www.boldsystems.org/) and GenBank (www.ncbi.nlm.nih.gov/nuccore/) to estimate the reliability with which a species identification can be obtained using DNA barcoding.

#### Material and Methods [1]



## Conclusion

Based on the present evaluation of the available sequence data, atpB, rps4 and trnL are the most promising DNA markers for the identification of *S. molesta*. However, due to the large gaps in available sequence data, it is currently impossible to fully assess the reliability of these markers.

## Discussion

DNA markers for which *Salvinia* sequences were available, were downloaded from GenBank and BOLD for all represented species of the genus *Salvinia* (as defined by [2]). Eight DNA markers were evaluated (Table 1). The species of priority to add in the dataset, since they belong a species complex, are: *S. auriculata, S. biloba, S. herzogii*. However, these were not represented in the online reference databases.

For markers **atpB**, **rps4** and **trnL**, the represented species (Table 2) have at least two sequences and clustering is supported for all. However, many and especially the most closely related species are not yet represented. Additional sequence data from *S. molesta*, covering both invasive and native range is needed as well. Hence, it is premature to decide about the ability of these DNA markers to differentiate *S. molesta* from other *Salvinia* species

For DNA marker **rbcL**, not enough sequence data is available. Two sequences of *Salviania molesta* cluster, but with little variation from a potentially mislabelled *S. cucullate*. Other species have only one sequence available.

For the intergenic spacer **psbA-trnH**, five sequences from the same study do cluster together, but two other sequence of *S. molesta* are not related and instead clustered with *S. minima*. Few congener are represented.

The genus is entirely underrepresented for markers **matK**, **ITS** and **trnG**. Therefore, it is currently impossible to assess the ability of these markers to identify *S. molesta*.

Table 1: Overview of	of the encour	ntered issues	concerning the	e DNA-based	identification	of the IAS [1	L]: (1) Insuffic	ient publicly	
available DNA seque	nces of the IA	AS to capture	the intra-specie	es divergence	; (2) Poor geo	graphical cove	rage of the IA	S sequences	
(native or invasive r	ange missing	;); (3) The IA	S sequences do	not form su	pported clust	ers; (4) Poten	itial misidenti	fication of a	
specimen which influ	lences the clu	ustering of th	e IAS sequences	; and (5) Not	all congenerio	c species are r	epresented in	the final NJ-	
tree. An 'X' indicate									
misidentified or mis						-	-		
applicable.	·		,		·				
Markers analysed	1		2		3			5	
rbcL	Х		Х			Х		Х	
matK	Х		X n/		/a			Х	
psbA-trnH	Х		X		X) X		Х		
ITS	Х		X 1		1			Х	
atpB	Х		Х				Х		
rps4	Х		Х					Х	
trnL	Х		Х					Х	
trnG	Х		Х					Х	
Table 2: Publicly ava	ilable sequen	ices downloa	ded (January 20	022) from BO	LD and GenBa	ank (including	sequences ex	tracted from	
plastid genomes) wh	ich were witl	hheld as relia	ble and information	ative in the fi	nal alignment	that was used	d for building	the NJ-trees.	
The species names for	ollow [2]. An '	'X' indicates	that at least one	e sequence w	as used in the	final alignme	nt, a '1' indica	ites only one	
sequences was availa	able for the fi	nal alignmen	t.						
Species in genus	rbcL	matK	psbA-trnH	ITS	atpB	rps4	trnL	trnG	
S. auriculata									
S. biloba									
S. cucullata	Х	Х			Х	Х			
S. hastata									
S. herzogii									
S. martynii									
S. minima	Х		Х	Х	Х	Х	Х	X	
S. minima <b>S. molesta</b>	X X	X	X X	X 1	X X	X X	X X	X X	
		X							
S. molesta	X	X	X	1	Х	X	X		
S. molesta S. natans	X	X	X	1	Х	X	X		
S. molesta S. natans S. nuriana S. nymphellula S. oblongifolia	X	X	X	1	Х	X	X		
S. molesta S. natans S. nuriana S. nymphellula	X X		X	1 X	X X	X X	X X	X	
S. molesta S. natans S. nuriana S. nymphellula S. oblongifolia	X X	X 2 /10-15	X	1 X	X X	X X	X X	X	

For a more elaborate discussion of the available databases, the sequence selection process, the outcome of the NJ-tree analyses, the usefulness of the investigated DNA sequences for species identification, as well as information on how to send samples for analyses please contact BopCo directly.

## **References and online information**

### Online information

https://nas.er.usgs.gov/queries/FactSheet.aspx?speciesID=298 https://www.daf.qld.gov.au/\_\_data/assets/pdf\_file/0003/65964/Salvinia-PP12.pdf http://www.gri.msstate.edu/research/invspec/factsheets/4P/Giant\_salvinia.pdf https://www.aphis.usda.gov/plant\_health/plant\_pest\_info/weeds/downloads/gsalvinia.pdf http://www.iucngisd.org/gisd/species.php?sc=569 https://www.fws.gov/fisheries/ans/erss/highrisk/ERSS-Salvinia-molesta-FINAL.pdf https://keys.lucidcentral.org/keys/v3/eafrinet/weeds/key/weeds/Media/Html/Salvinia\_molesta\_(Kariba\_Weed).htm http://especes-exotiques-envahissantes.fr/espece/salvinia-molesta/[FR]

#### **Picture credits**

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Page 2 (left, bottom): Salvinia (Salvinia molesta) is one of the world's worst aquatic weeds BY CSIRO [CC BY 3.0]

Page 2 (right): Salvinia molesta By John Forlonge [CC BY-NC-ND 2.0]

#### References

- [1] N. Smitz, S. Gombeer, K. Meganck, A. Vanderheyden, Y. R. Van Bourgonie, T. Backeljau, and M. De Meyer, "Identifying IAS based on DNA barcoding using currently available sequence data: details on applied material and methods." 2019. [Online]. Available from: http://bopco.myspecies.info/content/invasive-alien-species-ias-factsheets
- [2] Y. Roskov *et al.*, "Species 2000 & ITIS Catalogue of Life, 2019 Annual Checklist" *Species 2000*, 2019. [Online]. Available: www.catalogueoflife.org/annual-checklist/2019.
- [3] N. S. Nagalingum, M. D. Nowak, and K. M. Pryer, "Assessing phylogenetic relationships in extant heterosporous ferns (Salviniales), with a focus on *Pilularia* and *Salvinia*" *Bot. J. Linn. Soc.*, vol. 157, no. 4, pp. 673–685, 2008.
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- [6] K. Parys and A. Mikulyuk, "Salvinia molesta (kariba weed)" CABI Invasive Species Compendium, 2019. [Online]. Available: https://www.cabi.org/isc/datasheet/48447. [Accessed: 26-Mar-2020].
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