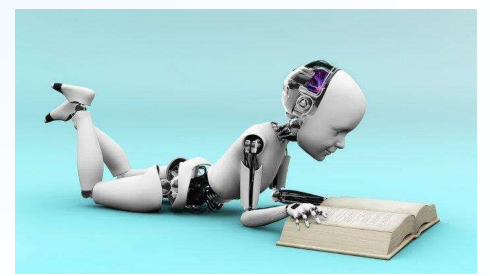
### 微生物组—扩增子16S分析研讨会第19期





# 32机器学习分类与回归

易生信 2023年6月11日



# 题 纲



○ 什么是机器学习?

o 随机森林和Adaboost的基本思想

o 两篇Nature结果解读

o RandomForest、Adaboost分析实战





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## 机器学习



 机器学习(Machine Learning, ML)是一门多领域交叉学科,涉及概率论、统计学、 逼近论、凸分析、算法复杂度理论等多门学科。专门研究计算机怎样模拟或实现 人类的学习行为,以获取新的知识或技能,重新组织已有的知识结构使之不断改 善自身的性能。



#### Nature近年来最高引的3篇文章全是深度学习

_	
Doon	learning.
Deen	i <del>c</del> ai i iii iu.

Y LeCun, Y Bengio, G Hinton Nature 521 (7553), 436-444

#### Human-level control through deep reinforcement learning.

V Mnih, K Kavukcuoglu, D Silver, AA Rusu, J Veness, MG Bellemare, ... Nature 518 (7540), 529-533

#### Mastering the game of Go with deep neural networks and tree search.

D Silver, A Huang, CJ Maddison, A Guez, L Sifre, J Schrittwieser, ... Nature 529 (7587), 484-489

#### Mastering the game of go without human knowledge



6101

5212

## 未来最热门的方法——深度学习



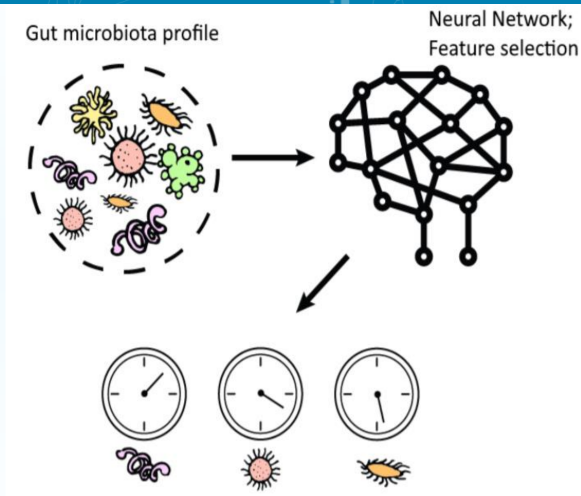
众所周知,目前最火的研究领域,莫过于 **肠道菌群和深度学习**,而此项研究的作者, 完美的将这两项研究结合,为我们带来一 项非常有趣的研究。该研究通过**深度学习** 的算法分析肠道菌群组成来预测人类年龄, 误差在4岁以内,说明菌群可作为生物标 志物用于衰老相关的研究。



Bacteroides are the most common bacteria species found in the human intestinal tract. DENNIS KUNKEL MICROSCOPY/SCIENCE SOURCE

#### The bacteria in your gut may reveal your true age

By Emily Mullin | Jan. 11, 2019, 11:50 AM



#### Effects bacteria elicit on age prediction

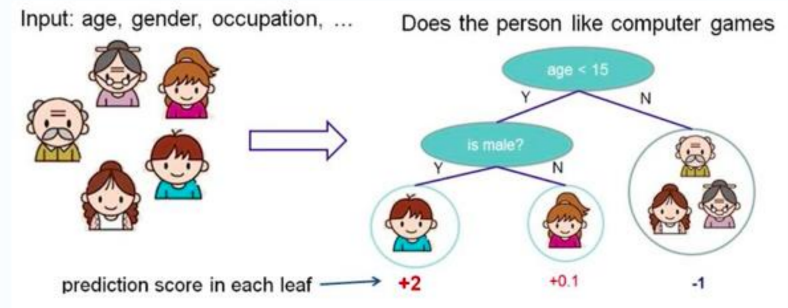


Science评论: 肠道菌群又添一大功能,揭示你的真实年龄 Fedor Galkin, Polina Mamoshina, Alex Aliper, Evgeny Putin, Vladimir Moskalev, Vadim N. Gladyshev, Alex Zhavoronkov. 2020. Human Gut Microbiome Aging Clock Based on Taxonomic Profiling and Deep Learning. *iScience* 23: 101199. https://doi.org/10.1016/j.isci.2020.101199 Nature Medicine同期8篇论文,聚焦人工智能在医学领域的应用

## 机器学习:十大常用算法



决策树、随机森林、逻辑回归、SVM、朴素贝叶斯、K最近邻、K均值、Adaboost、神经网络、马尔可夫



决策树:根据一些特征(feature)进行分类,每个节点提一个问题,通过判断,将数据分为两类,再继续提问。这些问题是根据已有数据学习出来的,再投入新数据的时候,就可以根据这棵树上的问题,将数据划分到合适的叶子上。



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## 随机森林

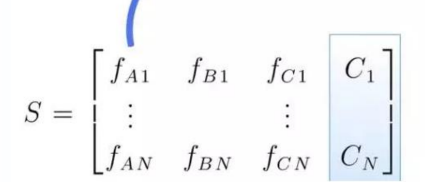


- 随机森林是一种基于决策树的高效的机器学习算法,可以用于对样本进行分类或回归分析。它属于非线性分类器,因此可以挖掘变量之间复杂的非线性的相互依赖关系。通过随机森林分析,可以找出能够区分两组样本间差异的关键成分(ASV/OTU/属/科等物种分类单元)。
- 优点:准确率极好;高效地分析大数据集上;处理高维特征的输入样本;缺省值结果很好;无需对它进行交叉验证或者用一个独立的测试集来获得误差的一个无偏估计(它可以在内部进行评估,也就是说在生成的过程中就可以对误差建立一个无偏估计)。
- o 注: 高水平文章为什么还要二次、三次验证呢?



## 随机森林的原理





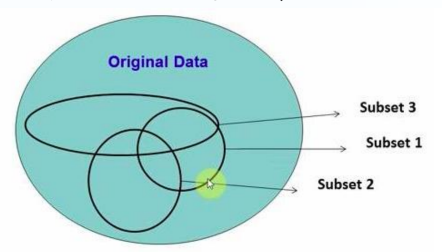
由 S 随机生成 M 个子矩阵 Create random subsets

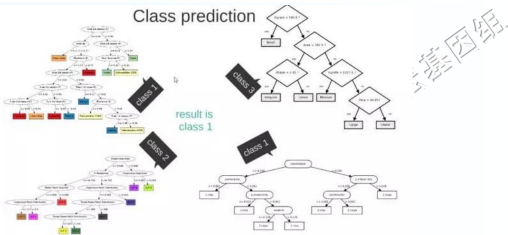
$$S_{1} = \begin{bmatrix} f_{A12} & f_{B12} & f_{C12} & C_{12} \\ f_{A15} & f_{B15} & f_{C15} & C_{15} \\ \vdots & & \vdots & & \vdots \\ f_{A35} & f_{B35} & f_{C35} & C_{35} \end{bmatrix} S_{2} = \begin{bmatrix} f_{A2} & f_{B2} & f_{C2} & C_{2} \\ f_{A6} & f_{B6} & f_{C6} & C_{6} \\ \vdots & & \vdots & & \vdots \\ f_{A20} & f_{B20} & f_{C20} & C_{20} \end{bmatrix}$$

$$\begin{array}{c} \text{Decision} \\ \text{tree 1} \\ S_{M} = \begin{bmatrix} f_{A4} & f_{B4} & f_{C4} & C_{4} \\ f_{A9} & f_{B9} & f_{C9} & C_{9} \\ \vdots & & \vdots & & \vdots \\ f_{A12} & f_{B12} & f_{C12} & C_{12} \end{bmatrix} \quad \begin{array}{c} \text{Decision} \\ \text{tree 2} \end{array}$$

$$\begin{array}{c} \text{Decision} \\ \text{tree M} \end{array}$$

在源数据中随机选取数据,组成M个子集



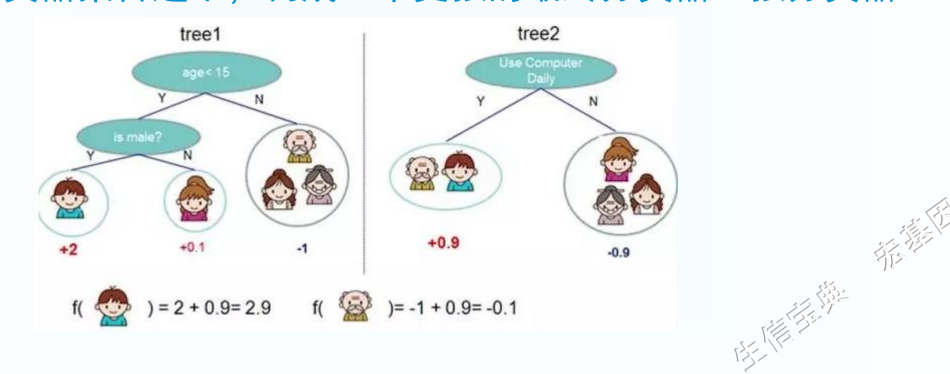


M 个子集得到 M 个决策树, 将新数据投入得到 M 个分类结果寻找哪些特征决策与己知分组一致率最高, 即分组贡献重要特征

## Adaboost ——迭代算法的一种



核心思想是针对同一个训练集训练不同的分类器(弱分类器),然后把 这些弱分类器集合起来,构成一个更强的最终分类器(强分类器)。



左右两个决策树,单个看是效果不怎么好的,但是把同样的数据投入进去, 把两个结果加起来考虑,就会增加可信度



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135 135 135



# ARTICLE



# Salt-responsive gut commensal modulates T<sub>H</sub>17 axis and disease

Nicola Wilck<sup>1,2,3,4,5</sup>, Mariana G. Matus<sup>6,7</sup>, Sean M. Kearney<sup>6</sup>, Scott W. Olesen<sup>6</sup>, Kristoffer Forslund<sup>8</sup>, Hendrik Bartolomaeus<sup>1,2,3,4</sup>, Stefanie Haase<sup>9</sup>, Anja Mähler<sup>1,5</sup>, András Balogh<sup>1,2,3,4,5</sup>, Lajos Markó<sup>1,2,3,4,5</sup>, Olga Vvedenskaya<sup>3,10,11</sup>, Friedrich H. Kleiner<sup>1</sup>, Dmitry Tsvetkov<sup>1,2</sup>, Lars Klug<sup>1,5</sup>, Paul I. Costea<sup>8</sup>, Shinichi Sunagawa<sup>8,12</sup>, Lisa Maier<sup>13</sup>, Natalia Rakova<sup>1,9</sup>, Valentin Schatz<sup>14</sup>, Patrick Neubert<sup>14</sup>, Christian Frätzer<sup>15</sup>, Alexander Krannich<sup>5</sup>, Maik Gollasch<sup>1,2,3</sup>, Diana A. Grohme<sup>16</sup>, Beatriz F. Côrte-Real<sup>17</sup>, Roman G. Gerlach<sup>18</sup>, Marijana Basic<sup>19</sup>, Athanasios Typas<sup>13</sup>, Chuan Wu<sup>20</sup>, Jens M. Titze<sup>21</sup>, Jonathan Jantsch<sup>14</sup>, Michael Boschmann<sup>1,5</sup>, Ralf Dechend<sup>1,2,5</sup>, Markus Kleinewietfeld<sup>16,17,22</sup>, Stefan Kempa<sup>3,5,10</sup>, Peer Bork<sup>3,8,23,24</sup>, Ralf A. Linker<sup>9</sup>§, Eric J. Alm<sup>6</sup>§ &









# 预测人高盐/低盐组生物标记(biomarker)+动物验证



a	Classifier	Max. relative	b	Animal	1	2	3	4	5	6	7	8	9	10	11
OTU	importance (%)	abundance (%)		Day –2											
Lactobacillus	25.0	1.24		Z Day -1											
Prevotellaceae	20.1	0.04		Dov. 1											
Pseudoflavonifract	or \ 12.5	0.85		Day 1											
Clostridia	12.5	0.58		Day 2											
Parasuterella	\ 9.7	7.33		O Day 2					$\sqsubseteq$	$\sqsubseteq$					
Akkermansia	9.3	4.22		$\perp$ Day 3											
Bacteroidetes	6.0	0.37		Day 14											
Alistipes	4.9	19.5		Day 14											
·				Class	ifica	ation	n pr	edi	ctic	n a	s [	$\square$ N	SD		] H

- 随机森林和Adabooster分类, Adabooster结果重要性importance值更大, 且对动物验证数据结果准确性也很高
- o 软件:使用Python3的slime2计算决定分组特征的贡献率/重要性

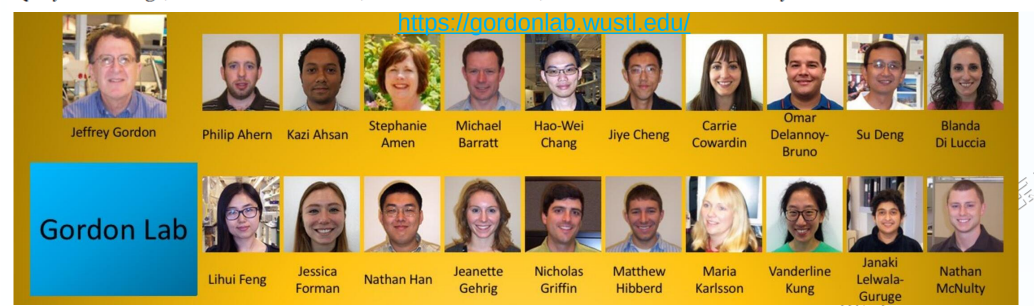


# LETTER



# Persistent gut microbiota immaturity in malnourished Bangladeshi children

Sathish Subramanian<sup>1</sup>, Sayeeda Huq<sup>2</sup>, Tanya Yatsunenko<sup>1</sup>, Rashidul Haque<sup>2</sup>, Mustafa Mahfuz<sup>2</sup>, Mohammed A. Alam<sup>2</sup>, Amber Benezra<sup>1,3</sup>, Joseph DeStefano<sup>1</sup>, Martin F. Meier<sup>1</sup>, Brian D. Muegge<sup>1</sup>, Michael J. Barratt<sup>1</sup>, Laura G. VanArendonk<sup>1</sup>, Qunyuan Zhang<sup>4</sup>, Michael A. Province<sup>4</sup>, William A. Petri Jr<sup>5</sup>, Tahmeed Ahmed<sup>2</sup> & Jeffrey I. Gordon<sup>1</sup>

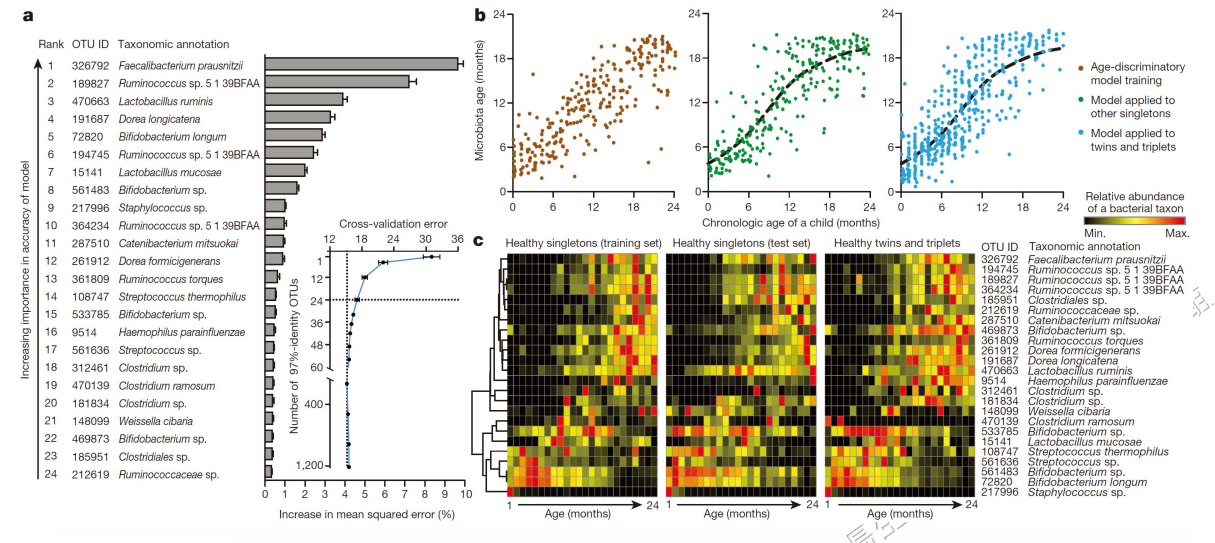




下次诺贝尔奖会是他吗? 肠道微生物组领域开创者Jeffrey Gordon Nature: 甘露糖苷选择性抑制致病性大肠杆菌 科普纪录片: 肠道菌群在人体中的作用

### 定义健康人1-24月菌群生物标记(Bacterial taxonomic biomarkers)







Subramanian et al. 2014. Persistent gut microbiota immaturity in malnourished Bangladeshi children. *Nature* 510: 417. https://doi.org/10.1038/nature13421

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# 扩展学习: Slime2鉴定分组贡献特征



# 使用adabooster计算10000次(16.7s), 推荐千万次

#### slime2.py otutab.txt design.txt --normalize --tag ab\_e4 ab -n 10000

Root; Bacteria; Firmicutes; Bacilli; Lactobacillales; Lactobacillaceae; Lactobacillus 0.2331

Root; Bacteria; Bacteroidetes; Bacteroidia; Bacteroidales; Prevotellaceae 0.1661

Root; Bacteria; Firmicutes; Clostridia; Clostridiales; Ruminococcaceae; Pseudoflavonifractor 0.1154

Root;Bacteria;Bacteroidetes 0.1071

Root; Bacteria; Firmicutes; Clostridia 0.0859

Root; Bacteria; Verrucomicrobia; Verrucomicrobiae; Verrucomicrobiales; Verrucomicrobiaceae; Akkermansia 0.0701

#### # 使用RandomForest计算10000次(14.5s),推荐百万次,支持多线程

#### slime2.py otutab.txt design.txt --normalize --tag rf\_e4 rf -n 10000

Root; Bacteria; Firmicutes; Bacilli; Lactobacillales; Lactobacillaceae; Lactobacillus 0.08616193880686196

Root;Bacteria;Proteobacteria;Betaproteobacteria;Burkholderiales;Sutterellaceae;Parasutterella 0.07444763171040213

Root; Bacteria; Verrucomicrobia; Verrucomicrobiae; Verrucomicrobiales; Verrucomicrobiaceae; Akkermansia 0.06552857763500938

Root; Bacteria; Firmicutes; Clostridia; Clostridiales; Ruminococcaceae; Pseudoflavonifractor 0.04065535850958773

Root; Bacteria; Firmicutes; Clostridia; Clostridiales; Ruminococcaceae; Oscillibacter 0.031399910016908436

Root; Bacteria; Firmicutes; Clostridia; Clostridiales; Lachnospiraceae 0.02733110442598093



# 分析需要OTU表和实验设计(分组信息)两个文件(演乐运行)

# R语言中的RandomForest包



#### [PDF] Classification and regression by randomForest

A Liaw, M Wiener - R news, 2002 - cogns.northwestern.edu

Recently there has been a lot of interest in "ensemble learning"—methods that generate many classifiers and aggregate their results. Two well-known methods are boosting (see, eg, Shapire et al., 1998) and bagging Breiman (1996) of classification trees. In boosting, successive trees give extra weight to points incorrectly predicted by earlier predictors. In the end, a weighted vote is taken for prediction. In bagging, successive trees do not depend on earlier trees—each is independently constructed using a bootstrap sample of the data set. In  $\ref{DD}$  Cite Cited by 20702 Related articles All 12 versions  $\ref{DD}$ 

- o 可以实现基于随机森林的分类(Classification)和回归(Regression)
- #安装install.packages("randomForest")
- o 查看帮助 ?randomForest



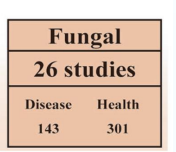


# 随机森林分类预测枯萎病的发生

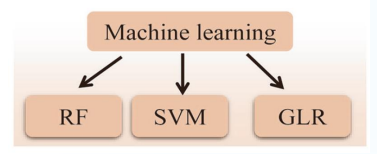




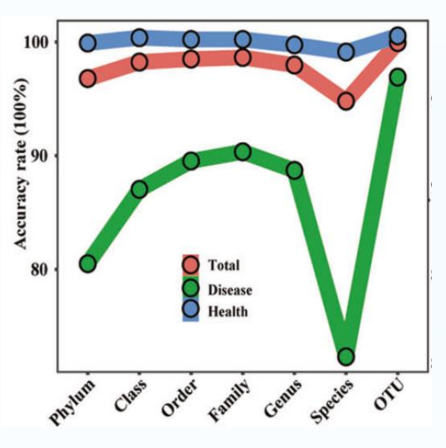
Bact	erial
37 stu	ıdies
Disease	Health
148	957

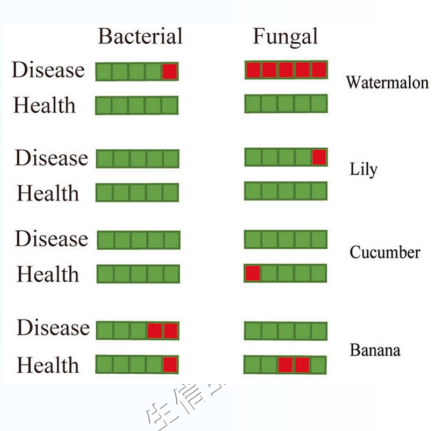


B



随机森林(RF)、支持向量机 (SVM)和逻辑回归(LR) 数据和方法选择





层级选择——分类预测评估准确度

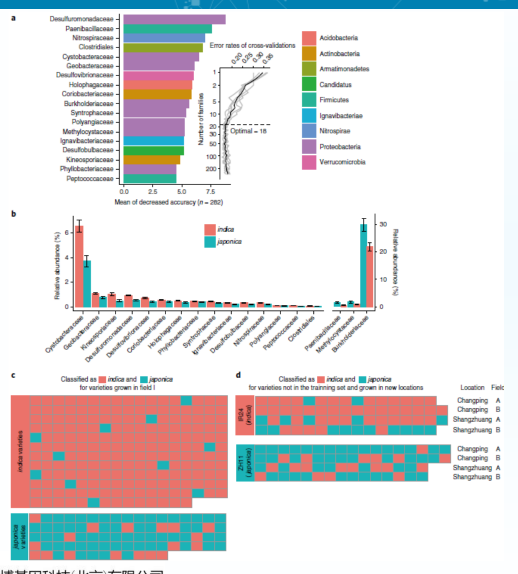
验证模型的有效性



ISME: 南农沈其荣团队基于大数据准确预测土壤的枯萎病发生 纯生信发 https://github.com/taowenmicro/Wen-etal-200214-paper-code 南京袁军/文涛ISME: 基于大数据整合准确预测土壤的枯萎病发生(视频)

# 随机森林分类实战: NBT基于微生物预测水稻亚种









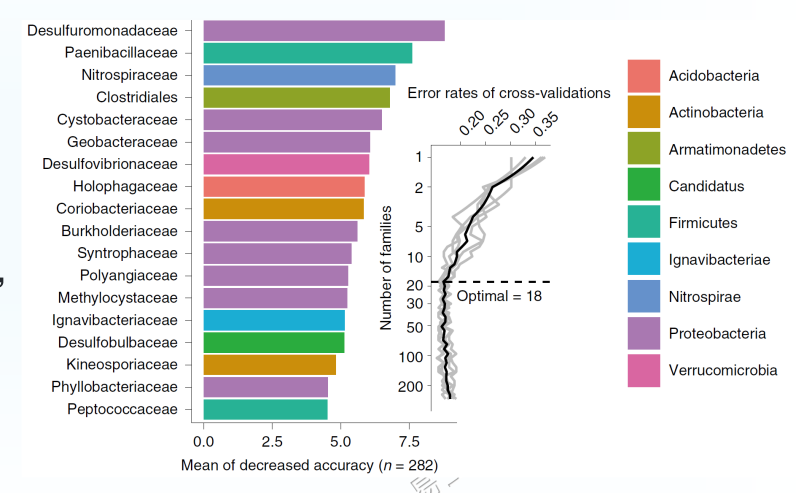
<sup>•</sup>NBT封面:水稻NRT1.1B基因调控根系微生物组参与氮利用

## 挖掘重要分类特征



randomForest包对样本 离散型进行分类,连续 型进行回归

- 1. randomForest建模和导出特征贡献度importance, ggplot2柱状图可视化
- 2. 多次交叉验证,人为选择 合适的特征组合, ggplot2折线图可视化

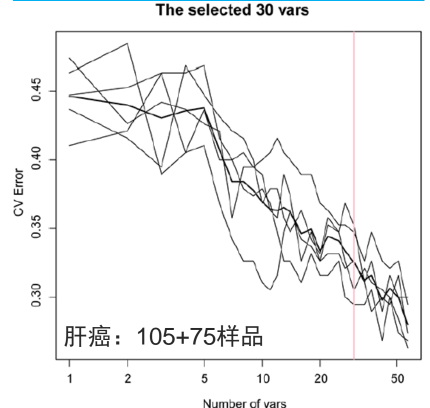




# 寻找生物标志物:交叉验证确定较优数量

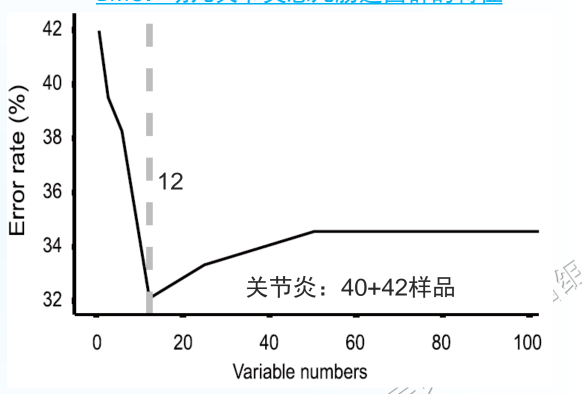


#### Gut-2018-早期肝癌肠道生物标志物鉴定



Zhigang Ren, Ang Li, Jianwen Jiang, Lin Zhou, Zujiang Yu, Haifeng Lu, Haiyang Xie, Xiaolong Chen, Li Shao, Ruiqing Zhang, Shaoyan Xu, Hua Zhang, Guangying Cui, Xinhua Chen, Ranran Sun, Hao Wen, Jan P. Lerut, Quancheng Kan, Lanjuan Li & Shusen Zheng. (2019). Gut microbiome analysis as a tool towards targeted non-invasive biomarkers for early hepatocellular carcinoma. *Gut* 68, 1014-1023, doi: https://doi.org/10.1136/gutjnl-2017-315084

BMC: 幼儿关节炎患儿肠道菌群的特征

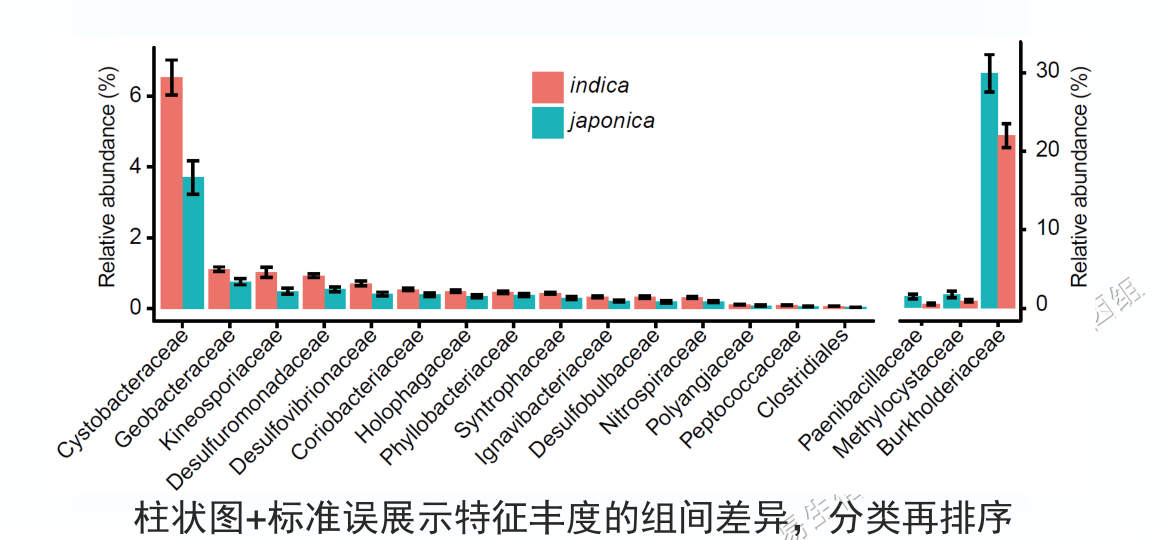


Xubo Qian<sup>†</sup>, **Yong-Xin Liu**<sup>†</sup>, Xiaohong Ye, Wenjie Zheng, Shaoxia Lv, Miaojun Mo, Jinjing Lin, Wenqin Wang, Weihan Wang, Xianning Zhang & Meiping Lu. (**2020**). Gut microbiota in children with juvenile idiopathic arthritis: characteristics, biomarker identification, and usefulness in clinical prediction. *BMC Genomics* 21, 286, doi: https://doi.org/10.1186/s12864-020-6703-0

易汉博基因科技(北京)有限公司 EHBIO Gene Technology (Beijing) co., LTD

## 重要分类特征在组间差异

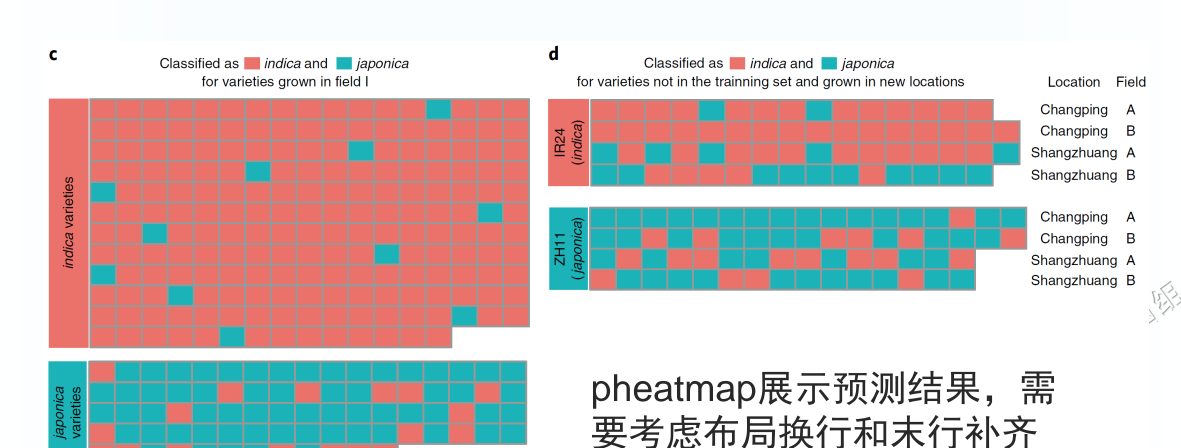






## 模型普适性评估——预测样本分类





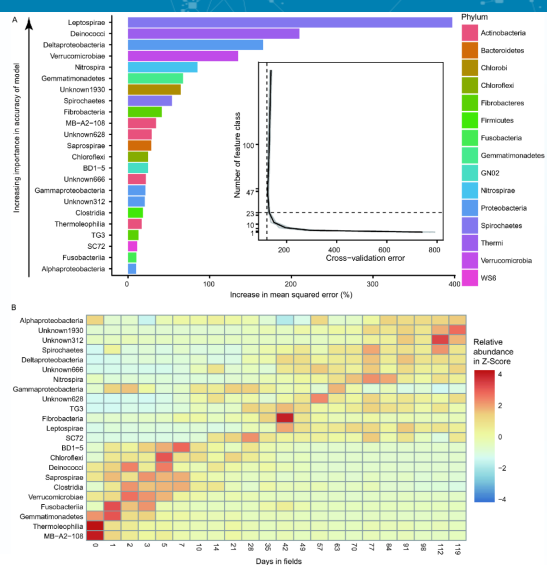
模型对另一块地相同品种(c)和不同地点不同品种(d)预测

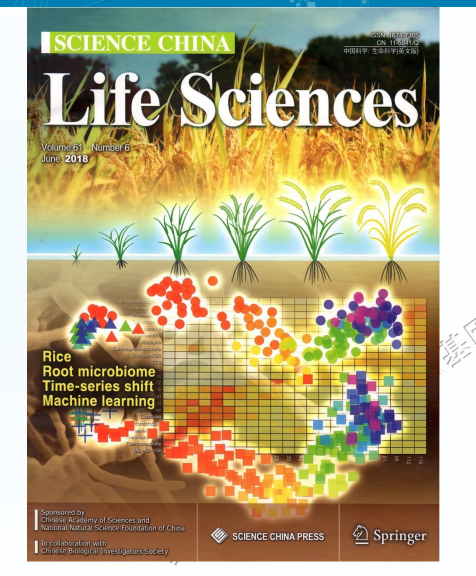


操作详见: RF\_classification目录

## R包RandomForest包分析回归实战









# 开始前准备



- o randomForest包
- o 实验设计: design.txt

SampleID	groupID	genotype	site	day	replicate	compartment
A50Cp0r1	A50Cp0	A50	Ср	0	1	root
A50Cp0r2	A50Cp0	A50	Ср	0	2	root
A50Cp0r3	A50Cp0	A50	Ср	0	3	root
IR24Cp0r1	IR24Cp0	IR24	Ср	0	1	root

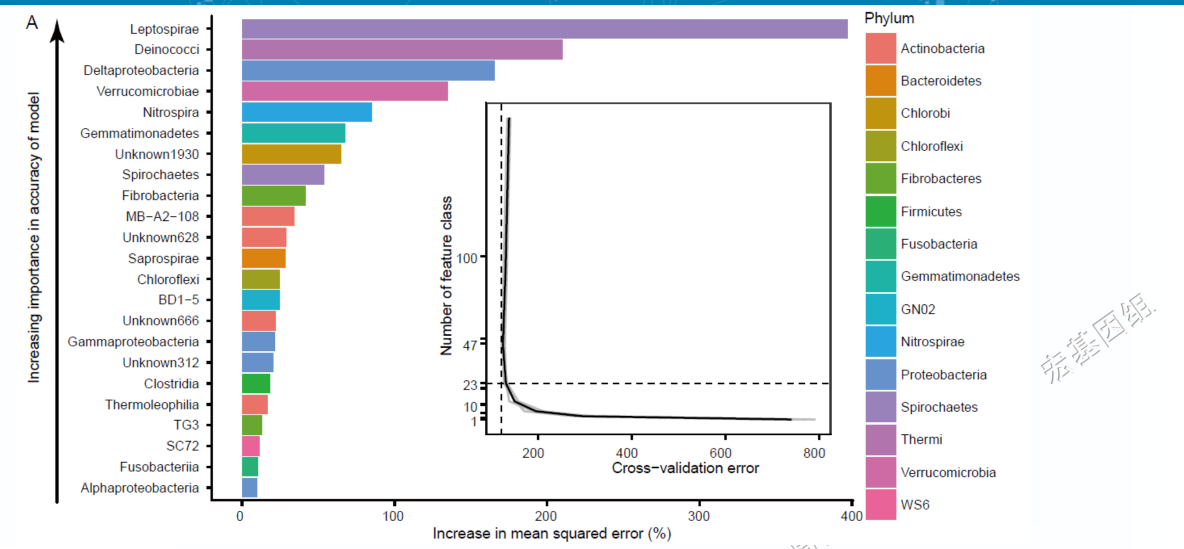
o OTU表/分类级: otu\_table\_tax\_L3.txt

ID The state of th	A50Cp10r2	A50Cp3r3	IR24Cp2r1
kBacteria;pProteobacteria;cBetaproteobacteria	0.29122	0.396278	0.316639
kBacteria;pProteobacteria;cAlphaproteobacteria	0.204275	0.162716	0.158932
kBacteria;pActinobacteria;cActinobacteria	0.218686	0.04667	0.12337
kBacteria;pProteobacteria;cDeltaproteobacteria	0.035686	0.04625	0.046986
kBacteria;pProteobacteria;cGammaproteobacteria	0.078237	0.159094	0.211824



# 水稻年龄(时间序列)相关Biomarkers挖掘



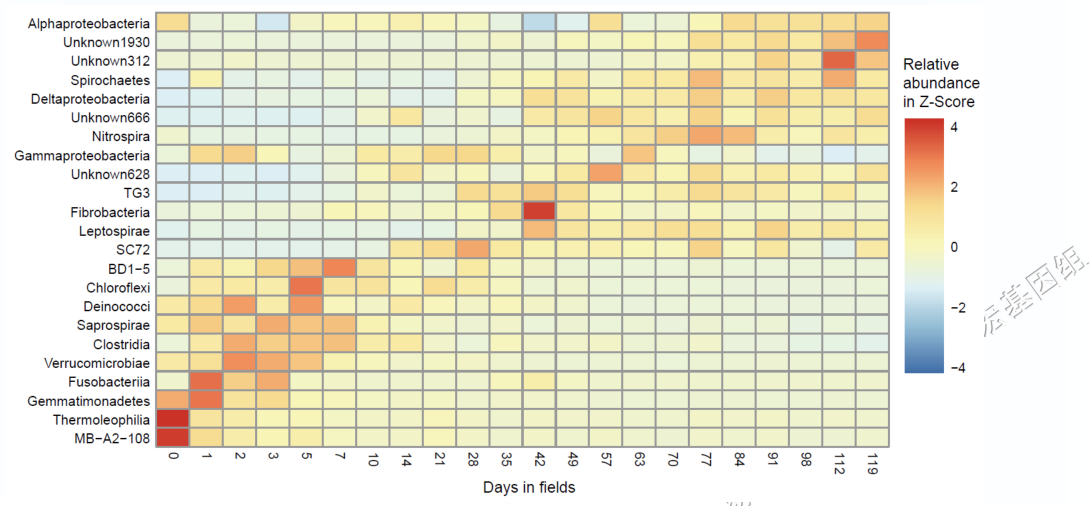




Jingying, Zhang<sup>(#)</sup>; Na, Zhang<sup>(#)</sup>; **Yong-Xin, Liu**<sup>(#)</sup>; Xiaoning, Zhang; Bin, Hu; Yuan, Qin; Haoran, Xu; Hui, Wang; Xiaoxuan, Guo; Jingmei, Qian; Wei, Wang; Pengfan, Zhang; Tao, Jin; Chengcai, Chu<sup>(\*)</sup>; Yang, Bai<sup>(\*)</sup>, Root microbiota shift in rice correlates with resident time in the field and developmental stage[J]. **Science China Life Sciences**, 2018: 1-9.

# 生物标志物在时间梯度上分布特征



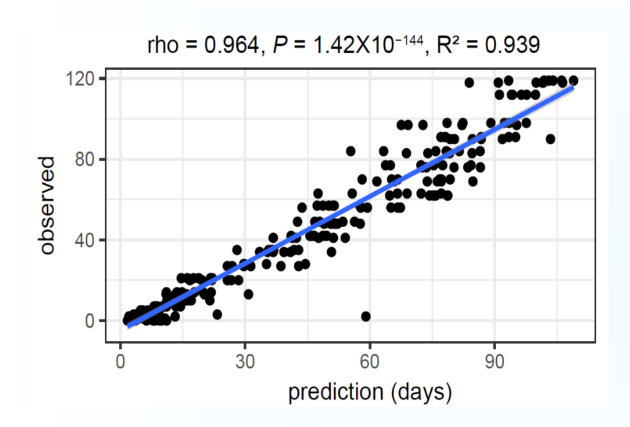


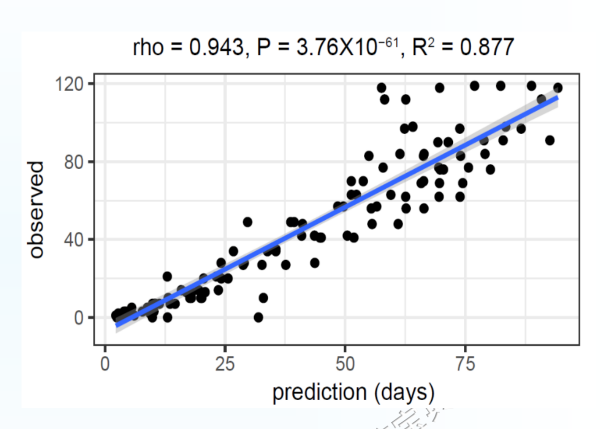


Liu et al. 2023. EasyAmplicon: An easy-to-use, open-source, reproducible, and community-based pipeline for amplicon data analysis in microbiome research. *iMeta* 2: e83. <a href="https://doi.org/10.1002/imt2.83">https://doi.org/10.1002/imt2.83</a>

# 基于细菌标志物模型预测水稻生育时期







训练集模型预测相关性

测试集模型预测相关性

Liu et al. 2023. EasyAmplicon: An easy-to-use, open-source, reproducible, and community-based pipeline for amplicon data analysis in microbiome research. *iMeta* 2: e83. <a href="https://doi.org/10.1002/imt2.83">https://doi.org/10.1002/imt2.83</a>



## 总结



- 机器学习领域是当前计算方法中热门的方法,其中深度学习将成本未来的研究和应用热点;
- 本领域目前已经开始应用深度学习,但使用最广泛的仍然是随机森林, 效果不佳时可尝试Adaboost;
- o Slime2可以实现随机森林和Adaboost分析,可在Linux下运行;
- R语言的RandomForest包引用超万次,可以Win/Mac/Linux跨平台分析,可实现随机森林分类与回归,同时结果R语言的统计和绘图完成全套的分析、统计与可视化——推荐。
- o QIIME 2: 15样品分类和回归q2-sample-classifier。







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