

Personalized prognosis & treatment using Ledley-Jaynes machines: An example study on conversion from Mild Cognitive Impairment to Alzheimer's Disease

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ABSTRACT

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1 INTRODUCTION: PERSONALIZED PROGNOSIS, TREATMENT, AND COMPUTER ALGORITHMS

1.0 Prologue: Four unique patients

Meet Olivia, Ariel, Bianca, Curtis. These four persons don't know each other, but they have something in common: they all suffer from a mild form of cognitive impairment, and are afraid that their impairment will turn into Alzheimer's Disease within a couple of years. This is why each of them recently underwent a battery of clinical examinations. Today they received the results. Based on these results, on available clinical statistical data, and on other relevant information, their clinician will assess their risk of developing Alzheimer's Disease. Then, together with the patients and their relatives, the clinician will make a decision among four distinct preventive-treatment options. In these tasks, the clinician will be helped by a computer algorithm.

Besides a shared diagnosis of Mild Cognitive Impairment and associated worries, these patients have other things in common – but also some differences. Let's take Olivia as reference, and list the similarities and differences between her and the other three patients:

- Olivia and Ariel have identical clinical results and age. They would also incur similar benefits and losses from the four available treatment options. Ariel, however, comes from a different geographical region, which presents a higher rate of conversion from Mild Cognitive Impairment to Alzheimer's Disease. And unlike Olivia, Ariel comes from a family with a heavy history of Alzheimer's Disease. Because of this geographical and family background, the clinician judges, before seeing the clinical data, that there's a 65% probability that Ariel's cognitive impairment will convert to Alzheimer's Disease.
- Olivia and Bianca have identical clinical results and age; they also come from the same geographical region and have very similar family histories. In fact, we shall see that they have the same probability of developing Alzheimer's Disease. Bianca, however, suffers from several allergies and additional clinical conditions that render some of the treatment options slightly riskier for her.
- Olivia and Curtis have different results on all clinical examinations; Olivia is also more than 10 years older than Curtis. They otherwise come from the same geographical region, have very similar family histories, and would incur similar benefits or losses from the treatment options. Note that one clinical result for Curtis's (hippocampal volume) is missing.

Considering the similarities and differences among these patients, which of the four available treatments will be optimal for each of them? The clinician will find that, despite the many factors in common among our four patients – even despite Olivia's, Ariel's, and Bianca's identical clinical results, and Olivia's and Bianca's identical probability of conversion to Alzheimer's Disease – the optimal treatment for each patient is different from those for the other three.

1.1 Assistive computer algorithms: personalized input and output

In the example above, we said "in these tasks, the clinician will be helped by a computer algorithm". The need for such computational help is clear from the vast amount of clinical statistical data and the large

¹ These are purely fictive characters but with clinically realistic conditions; any reference to real persons is purely coincidental.

² In the present paper, we use "prognosis" in a general sense to include also "diagnosis", and "treatment" quite loosely to mean any course of action a clinician might take, including "preventive treatment", or even "additional tests".

number of clinical predictors today available to clinicians. But how should such an assistive computer algorithm be designed in order to take fully into account patient differences?

Although the example above concerns specifically Alzheimer's Disease, the differences among patients

described there apply more generally to most, if not all, clinical problems of prognosis and treatment. These differences can be broadly categorized as "difference in auxiliary and background information" (Olivia and Ariel), "difference in benefit and availability of treatments" (Olivia and Bianca), "difference in clinical predictors" (Olivia and Curtis), as schematized in the side figure. Each of these difference categories can affect the clinician's final choice of optimal treatment. An assistive algorithm should therefore reflect these differences in its input, its output, or both:



- In principle, there could be three kinds of input "slots", where the clinician can input the current patient's specific values as regards clinical predictors, auxiliary information, and treatment options & benefits.
- If input slots are only available for one or two of the categories above, the output should at least be of such a kind as to allow the clinician to integrate the current patient's specific values of the missing input categories.

To appreciate these requirements, contrast the input and output of many kinds of machine-learning classification algorithms. These typically only allow the input of a patient's clinical predictors, with no space for patient-specific auxiliary information or for adjustments of differences in background statistics (think of Olivia and Ariel). And they typically output only a discrete prognostic label (say, "stable Mild Cognitive Impairment" vs "conversion to Alzheimer's Disease"), but no measure of the uncertainty about that label. Unfortunately, such output does not allow the clinician to assess treatment benefits and losses for the current patient, for this assessment depends not on the presence (present or future) of a disease, but on the *risk* of its presence. We shall discuss these points at length in §§ 3.2 and 3.3.

The purpose of the present work is to present an assistive algorithm that meets the requirements above. This algorithm is designed to first learn from a dataset of clinical data with relevant predictors and predictand³, and then assist a clinician in the assessment of prognosis & treatment for new patients. It offers these ten features:

- 1. It can work with clinical predictors comprising any combination of categorical and one-dimensional (continuous, discrete ordinal, unbounded or bounded, uncensored or censored) variates. The predictand can also be any combination of categorical and one-dimensional variates.
- 2. It treats predictor and predictand variates on equal footing, in the sense that the clinician can at any moment decide to predict some other variate given the rest.
- 3. It does not require that the current patient be considered in all respects as a member of the population underlying the learning dataset. The patient can be considered a member only conditionally on particular variate values.
- 4. It accepts three inputs:
 - a. the clinical-predictor values for the current patient;

³ literally "quantity to be predicted" (cf. *measurand* in metrology, JCGM 2012, 2.3). We find this term, used in meteorology and climate science, more precise and less obscure or misleading than "dependent variate", "response variate", or similar.

- b. information about which predictand-predictor relationships learned from the dataset can be generalized to the current patient, and a prior prognostic probability representing auxiliary information;
- c. a set of treatment options and their benefits and losses for the current patient.
- 5. It yields three basic outputs
 - a. any prognostic probabilities or likelihoods about predictors and predictand desired by the clinician, given input 4a;
 - b. final prognostic probabilities, given inputs 4a–4b;
 - c. optimal treatment, given inputs 4a-4c;
- 6. Its input and outputs are modular, in the sense that the clinician can, for instance, give inputs 4a–4b only, get a prognostic probability 5b as output, and then proceed to treatment assessment by other means or algorithms.
- 7. It works even if predictor data are missing, both in the learning dataset and for the current patient.
- 8. It can quantify the uncertainty of its own outputs and therefore allow for sensitivity analyses. For example, it can tell how much a prognostic probability could have been different if the learning dataset had been larger, or whether the optimal treatment could be different if a particular missing predictor for the current patient had been available.
- 9. It can make various kinds of long-term forecasts, such as frequency of prognoses having given probabilities, frequency of prescribed treatments, and similar provided that the dataset used for its learning can be considered representative of the full population,
- 10. It is model-free and extracts the maximal amount of information theoretically contained in the learning dataset, and therefore achieves the maximal prognostic power that the predictors can yield. In other words, it is unbeatable.

Let us comment on some of these features. We believe that the capability of working with complex predictands, feature 1, is important for a more realistic and nuanced approach to prognosis. In the case of Alzheimer's Disease, for instance, it is clear that a simple dichotomy "has disease" vs. "doesn't have disease" is an oversimplification more text and references?. Without the capability of being used with patients having peculiar contexts or clinical situations, feature 3, the algorithm would be of no use in the often occurring case of patients having special clinical contexts. The capability of dealing with missing data, feature 7, is important for a concrete implementation in a clinical setting, typically afflicted by imputation problems. Feature 8 is extremely important for a clinician to assess the reliability of final decisions and inform the patient of the possibility of unwanted outcomes. Finally, features 2 and 10, the fact that this algorithm yields the maximal amount of information (as determined by information theory) jointly contained in all variates, makes it valuable in general clinical research. The algorithm can, for example, forecast the maximal accuracy obtainable by *any* inference algorithm based on the same predictors or another set of predictors; it itself attains that maximal accuracy.

Further features relevant for Machine Learning are given in § ***.

We call this algorithm a *Ledley-Jaynes machine*, for reasons explained in the next section. It is at the moment available as a collection of scripts⁴ in the R programming language (R Core Team, 2023), which we plan to soon assemble into a clinician-friendly R package.

⁴ https://github.com/pglpm/ledley-jaynes_machine

The next section 2 gives an intuitive understanding of the Ledley-Jaynes machine's underlying principles and workings. It is largely independent of the subsequent sections, so it can be skipped by readers who want to immediately see its application. Application is shown in § 3, using the four-patient fictitious scenario of § 1.0 as a concrete example; the final subsection 3.4 discusses applications to general medical research. A summary and discussion is given in § 4.

Mathematical details and proofs on which the present work is grounded are given in a companion technical note⁵, which explains more in detail how to use the R scripts.

2 THE LEDLEY-JAYNES MACHINE

This section can be skipped by readers not interested in the principles and mathematical details underlying the Ledley-Jaynes machine.

2.1 Underlying theory and characteristics

The method to solve clinical decision-making problems such as the one of § 1 is none other than Decision Theory: the combination of probability theory and utility theory. It integrates available clinical statistical data with each patient's unique combination of clinical results, auxiliary information, and treatment benefits, in a mathematical framework, completely determined by basic self-consistency requirements.⁶

Medicine has the distinction of having been one of the first fields to adopt Decision Theory, with the pioneering work by Ledley and Lusted (1959a; 1959b; 1960; 1960), who also promoted its algorithmic implementation (Ledley, 1959, 1960, see especially § 1-5 p. 21). Clinical decision-making is today explained and exemplified in brilliant textbooks for medical students and clinicians (Weinstein and Fineberg, 1980; Sox et al., 2013; Hunink et al., 2014). An outline is given in § 3.3.

The "Ledley-Jaynes machine" is an algorithmic implementation of the main calculations underlying the clinical decision-making process, as dreamed by Ledley and Lusted: from the comparison of a patient's specific predictors with the statistics offered by a clinical database, to the choice of optimal treatment. The name is a homage to Ledley – who, incidentally, died of Alzheimer's Disease (Shah et al., 2013) – and to Jaynes (2003), who brilliantly explained the inductive logic underlying such a "robot".

Decision theory is also the normative foundation for the construction of an Artificial Intelligence agent capable of rational inference and decision making (Russell and Norvig 2022, part IV; Jaynes 2003, chs 1–2, 13–14). The Ledley-Jaynes machine can therefore be seen as an *ideal* machine-learning algorithm. It is "ideal" in the sense of being free from special modelling assumptions (this is why we do not call it a "model") and from limitations of its informational output, characteristic of most common machine-learning algorithms; not "ideal" in the sense of being impracticable. Quite the opposite, the present work shows that this ideal machine-learning algorithm can today be applied at insubstantial computational cost.

More concretely, the Ledley-Jaynes machine is ideal because *it computes the probability distribution* over all possible long-run joint frequency distributions from which the learning dataset can originate (joint distributions for all predictor and predictand variates). This is the maximum possible amount of information that can be extracted from the learning dataset, in a strict information-theoretic sense. From

 $^{^{5} \ \}text{https://github.com/pglpm/ledley-jaynes_machine/raw/main/ledley-jaynes_machine.pdf}$

⁶ Jaynes (2003); von Neumann and Morgenstern (1955); Cox (1946); Savage (1972); Luce and Raiffa (1957); Raiffa and Schlaifer (2000); Raiffa (1970); Lindley (1988); Kreps (1988, chs 13–14).

this probability distribution, the Ledley-Jaynes machine can indeed calculate any quantity outputted by other machine-learning algorithms. For example (for terminology see e.g. Murphy, 2012, § 8.6):

- "Discriminative" algorithms: the probability $p(Y \mid X)$ of any set of predictands Y given any set of input predictors X.
- "Generative" algorithms: the probability $p(X \mid Y)$ of any set of input predictors X given any set of predictand values Y.
 - More generally, the machine can compute any joint, marginal, or conditional probabilities p(Z', Z''), p(Z'), $p(Z' \mid Z'')$ for any desired subsets of variates Z', Z''.
- Regression or classification: the average value of any set of variates Y, given any other set of variates X, including the special case of predictand Y and predictors X. The uncertainty or variability around such an average is also automatically computed.
- Functional regression: if the predictand Y or any other variate of interest turns out to be a function f of variates X, then their conditional probability will be a delta distribution: $p(Y \mid X) = \delta[Y f(X)]$. Thus the Ledley-Jaynes machine always recovers a functional relationship if there is one, as well as its noise distribution.

Furthermore, the machine also quantifies the uncertainty of all outputs above. More precisely, it takes into account that the statistical properties of the learning dataset could be different from those of its original population, owing to sampling fluctuations; and it computes how much any of the outputs above would probably change if more learning data were collected.

In the next section, we explain intuitively how the Ledley-Jaynes machine computes the general probability distribution over long-run frequencies; but we can already summarize a couple of special characteristics brought about by such computation. Unlike machine-learning algorithms such as neural networks, random forests, support-vector machines, logistic regression, or generalized linear models, the Ledley-Jaynes machine does not do an optimization during the learning phase, searching for the minimum of some objective function. It does a full hypothesis-space survey. The optimization done by most machine-learning algorithms is an approximate form of this survey, based on the assumption or hope that the most relevant portion of the hypothesis space will be around the extremum (MacKay, 1992a; Murphy, 2012, ch. 16; see also Self and Cheeseman, 1987). The underlying necessity of a more extensive survey, however, becomes manifest in many of the obligatory procedures that go together with the training of most machine-learning algorithms, cross-validation being a prominent example (MacKay, 1992b). This leads to another special characteristic: the Ledley-Jaynes machine does not need validation sets, test sets, or other data splits; nor does it need cross-validation procedures. Intuitively this is the case because the underlying hypothesis-space survey realizes a sort of full-fledged cross-validation and data partition. It can indeed be proved that one of the internal computations of the machine is mathematically equivalent to doing k-fold cross-validations for all possible data splits and k (Porta Mana, 2019; Fong and Holmes, 2020).

Such flexibility and informationally rich output come, of course, at a computational cost. Until some years ago, the cost would have been prohibitive in all but the simplest inferential problems. But today, an inference problem involving 13 variates and 700 datapoints, such as the example considered in the present work, takes less than six hours of computation on an office computer workstation. We discuss computational limitations further in § 4.2.

2.2 Intuitive understanding of the learning algorithm

The calculations by which the Ledley-Jaynes machine learns and operates are univocally determined by Cox's theorem ⁷, which determines quantitative inference rules from self-consistency requirements, and by de Finetti's theorem⁸, which further constrains these rules in the case of "generalization from similar cases". These calculations have a very intuitive interpretation.

We consider a patient to be a member of some population of similar past, present, and future patients. Suppose we knew the joint frequency distribution of all possible combinations of predictor and predictand values in such a population. We would then judge the probability for a patient's particular variate values to be equal to their corresponding population frequency. Pure symmetry considerations lead to this result (de Finetti, 1930; Dawid, 2013; Bernardo and Smith, 2000, §§ 4.2–4.3). The same would be true for conditional and marginal probabilities and frequencies. This population frequency distribution would bound the maximal prognostic power attainable with the given predictors in the population. A higher prognostic power could only be attainable by using additional or different predictors having sharper conditional frequencies for the predictand in the population. Given knowledge of such frequency distribution, there would be no problem of "generalizing" to new patients, because each new patient would already be counted in the known frequencies. An inference algorithm would only need to enumerate and memorize, rather than to learn and generalize.

Learning and generalization come into play because the frequency distribution for the population is unknown: we only have a sample from it, the "learning dataset". Thus we can, at most, assign a probability to each possible frequency distribution. This is precisely what the Ledley-Jaynes machine does.

The way in which the machine assigns a probability to each "candidate" true frequency distribution is also intuitive. It combines two factors: (i) how well the candidate fits the sample dataset, (ii) how biologically or physically reasonable the candidate is. The first factor is easily computed: it is the joint probability of the dataset if it were sampled from a population having that candidate frequency. The second factor is a prior probability expressing how reasonable that candidate is. ¹⁰ The most general natural requirement for "reasonableness" is that the candidate should have some degree of smoothness, owing to physical and biological reasons. Figure 1a shows samples of what the machine considers to be "reasonable candidates" for the population frequency distribution of a discrete variate. Note that no frequency distributions are excluded; they are only given higher or lower probabilities.

The product of the two factors (i), (ii), normalized, yields the probability of each possible frequency distribution. An illustration of factors (i), (ii) at work is given in fig. 1b for an example problem with two variates. The Ledley-Jaynes machine outputs the distribution of these final probabilities in the form of a large sample (the amount is decided by the user) drawn from it. In this form, all other marginal or conditional probabilities and averages of interest are calculated via Monte Carlo integration. This is the same methodology, but in nonparametric form, used in various inferences about the black holes M87 and Sagittarius A* (Event Horizon Telescope Collaboration, 2019, 2022).

⁷ Cox, 1946, 1961; Pólya, 1954, 1968; Tribus, 1969; Fine, 1973; Rosenkrantz, 1977; Paris, 2006; Snow, 1998; Arnborg and Sjödin, 2001; Snow, 2001; Clayton and Waddington, 2017; for a review see Van Horn, 2003.

⁸ de Finetti, 1930, 1937; Bernardo and Smith, 2000, §§ 4.2–4.3; for a review see Dawid, 2013.

⁹ If there were a functional relationship from predictors to predictand, then the predictand value corresponding to the function output would have conditional frequency and probability equal to 1, and all other values to 0. Therefore, this point of view still encompasses a functional relationship as a particular case.

¹⁰ It must be emphasized that some notion of "reasonable candidate" is unavoidable and clearly present in the construction or testing of any inference algorithm. How can we otherwise judge that an algorithm is over- or under-fitting? Such judgement implies that we have a preconceived, reasonable reference distribution in the back of our minds. The fit is either intuitively compared with the preconceived reference; or is compared with a known, ground-truth test for testing owing to its similarity with the preconceived reference.



Figure 1a. Samples of initially probable candidates of the true population frequency distribution, for a variate such as RAVLT-del or RAVLT-rec.

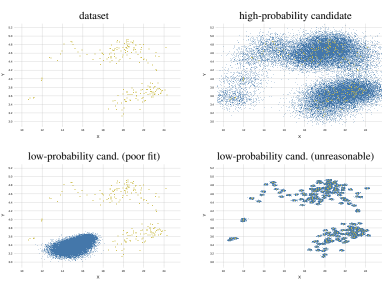


Figure 1b. Illustration of the two factors determining the final probability of a candidate population frequency distribution. *Upper-left*: an example dataset with two variates. *Upper-right*: candidate frequency distribution with high final probability; it fits the data and is reasonable. *Lower-left*: candidate distribution with low final probability; it is reasonable but does not fit the data. *Lower-right*: candidate distribution with low final probability; it fits the data but is not reasonable.

3 EXAMPLE APPLICATION

In this section, we illustrate how the Ledley-Jaynes machine is applied in the example case outlined in § 1.0. Although the patients are fictitious, the dataset is real and briefly discussed in the next subsection. The main inferential and decision-making steps are summarized in table 1. As mentioned in § 1.1, steps 1.–3. are modular: the clinician is free to stop after any of them and use their output in other ways or with other algorithms.

These steps are illustrated in the next three subsections, preceded by an explanation of their rationale. They are presented in chronological order as the clinician would apply them. Steps 1.–3. could also be presented in reverse order; which would be more suited to their logical dependence, as the procedure in each step is actually motivated by the one in the next. We suggest that readers familiar with the principles of clinical decision-making read the following subsections in 3.0–3.1–3.2–3.3 order; whereas readers unfamiliar with these principles read them in 3.0–3.3–3.2–3.1 order.

3.0 Predictors, predictand, and learning dataset

The dataset used in our example comes from the study by the Alzheimer's Disease Neuroimaging Initiative (ADNI). This longitudinal multicentre study is designed to develop and validate neuroimaging and biochemical biomarkers for the early detection, monitoring, and treatment of Alzheimer's Disease (Petersen et al., 2010). The present dataset consists of 704 ADNI subjects meeting the criteria for Mild Cognitive Impairment at their first, baseline assessment who additionally had a minimum of two following, additional study visits, as well as three MRI examinations. Each subject's diagnostic status was reevaluated at each study visit. This longitudinal diagnostic label is used as the predictand variate CAD in our study: it categorizes each subject as either converting to Alzheimer's Disease after the first study visit (CAD = 1) or

machine.

Table 1. Main inferential and decision-making steps

- Find or build an appropriate dataset of clinical cases comprising values of the predictors and predictand
 of interest. Datapoints with partially missing values are allowed.

 Input the dataset into the Ledley-Jaynes machine and let it infer the joint full-population frequencies of
 predictors and predictand behind the dataset.
- 1. Measure the present patient's predictor values and input them in the Ledley-Jaynes machine. Partially missing values are allowed.
- 2. Assess which conditional statistics of the dataset can be applied to the present patient, and any auxiliary clinical information available. Quantify the latter it in a prior probability.

 Input the relevant statistics and auxiliary information for the present patient into the Ledley-Jaynes
 - Upon request, the machine can now output the final probability of the predictand's true value for the patient, as well as any other probabilities and likelihoods of interest.
- 3. Assess the clinical courses of action (treatments, more tests, and so on) available for the present patient, and the utility (benefit and loss) of each course of action, depending on each possible predictand value for the patient.
 - Input the patient's utilities into the Ledley-Jaynes machine. The machine outputs the course of action having maximal expected utility.
 - Upon request, the machine can output the probability of gaining different utilities, perform sensitivity analyses for missing data, and do other similar tasks.

remaining stable with Mild Cognitive Impairment (CAD = 0). The dataset has 325 subjects (46.2%) with CAD = 1 and 379 (53.8%) with CAD = 0. Criteria used for classifying subjects as having Mild Cognitive Impairment or Alzheimer's Disease, as well as ADNI's general criteria for subject inclusion, are described in McKhann et al. (1984); Petersen et al. (2010).

The 12 predictor variates consist of the results of eight tests for cognitive function (ANART, CFT, GDS, RAVLT-imm, RAVLT-del, RAVLT-rec, TMTA, TMTB), the presence of the APOE4-e4 risk allele (Liu et al., 2013), a normalized measure of hippocampal volume (HV), Age, and Sex. Further details about these variates and their selection can be found in Rye et al. (2022). Add details about specially sampled variates (age?), important for step 2. The cognitive variates are integer-valued; hippocampal volume and age are continuous, and APOE4 and sex are binary. The values of one or two of these predictors were missing for 30 subjects in the dataset.

The Ledley-Jaynes machine took less than five hours (on a 16-core Intel Core i9-12900K CPU) to calculate the probability distribution for the possible population joint frequencies distributions of the 13 variates. Some results can already be visualized after this inference.

Figure 2 shows, on the left, the predicted distributions of RAVLT-del, RAVLT-imm, GDS, and hippocampal volume for the subpopulation of patients that will convert to Alzheimer's Disease (red dashed) and the subpopulation that will remain with stable Mild Cognitive Impairment (solid blue). On the right, the predicted frequency of conversion in the full population is given, conditional on the same predictors. The thin curves are 100 samples of highly probable frequency distributions; the thicker lines are their means, which are also the predictive conditional probabilities. The two subpopulations of patients are clearly distinct in the RAVLT-del, RAVLT-imm, HV variates. These predictors can yield probabilities of

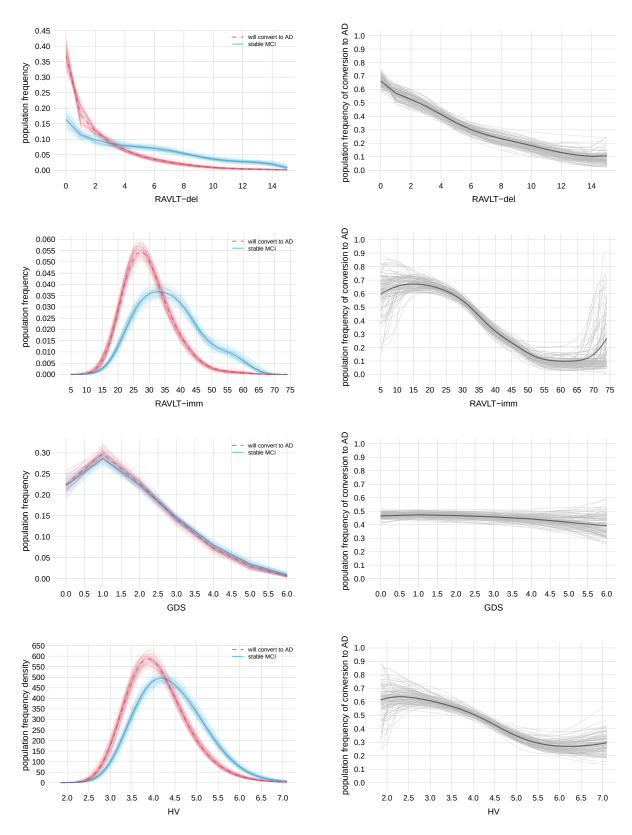


Figure 2. Predicted distributions of some predictor variates, for the subpopulation of patients that will convert to Alzheimer's Disease (red dashed) and the subpopulation with stable Mild Cognitive Impairment (solid blue).

conversion as high as 70% or as low as 10%. The two subpopulations are practically indistinguishable in the GDS variate, which, therefore, always gives very uncertain predictions.

The learning dataset comprises enough data to greatly reduce our uncertainty about the population distributions, as evident from the very narrow spread of the curves. These simple results show the great usefulness of the Ledley-Jaynes machine for general medical research.

3.1 Patient's clinical information

The 12 predictor values for our four patients are reported in table 2, top. Note that Curtis's value for the Hippocampal Volume is missing; this is not a problem for the Ledley-Jaynes machine. Given these predictor values the Ledley-Jaynes machine can output any probabilities of interest to the clinician. Table 2, bottom, reports three probabilities that are important for the step of the next subsection:¹¹

- p(cAD = Y | predictors): the probability that the patient will convert to Alzheimer's Disease, given the patient's specific predictors and that the patient comes from the same population as the learning dataset.
- $p(predictors \mid cAD = Y)$: the probability that a patient who will convert to Alzheimer's Disease would have these specific predictor values. In other words, the *likelihood*¹² of conversion to Alzheimer's Disease, given the predictors.
- p(predictors | CAD = N): the probability that a patient who will remain with stable Mild Cognitive Impairment would have these specific predictor values. In other words, the *likelihood* of stable Mild Cognitive Impairment, given the predictors.

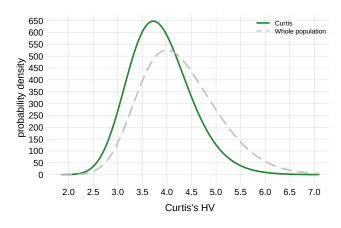
The Ledley-Jaynes machine can also answers other questions of interest to the clinician. For instance, what could be the value of Curtis's Hippocampal Volume? The answer is given in fig. 3a, which also

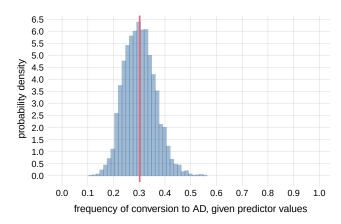
 $^{^{12}}$ p(A | B) is the probability of A given B, as well as the likelihood of B given A (Good, 1950, § 6.1).

	Olivia	Ariel	Bianca	Curtis
Age Sex	75.4 F	75.4 F	75.4 F	63.8 M
Hippocampal volume $(HV)/10^{-3}$	4.26	4.26	4.26	[missing]
APOE4 status	N	N	N	Y
Reading test (ANART)	18	18	18	15
Category Fluency Test (CFT)	21	21	21	14
Geriatric Depression Scale (GDS)	3	3	3	2
RAVLT-imm ediate memory	36	36	36	20
RAVLT-del ayed recall	5	5	5	0
RAVLT-recognition	10	10	10	3
Trail Making Test A (TMTA)	21	21	21	36
Trail Making Test B (TMTB)	114	114	114	126
$\begin{array}{c c} p(\texttt{cAD} = \texttt{Y} \mid \texttt{predictors}) \\ p(\texttt{predictors} \mid \texttt{cAD} = \texttt{Y})/10^{-12} \\ p(\texttt{predictors} \mid \texttt{cAD} = \texttt{N})/10^{-12} \end{array}$	0.302 8.97 18.6	0.302 8.97 18.6	0.302 8.97 18.6	0.703 1.14 0.343

Table 2. Predictor values for the four patients, and resulting conditional probabilities

All relative uncertainties of the results are below 0.8%, Curtis's two likelihoods being an exception at 2%.





pocampal Volume

Figure 3a. Probability distribution for Curtis's Hip- **Figure 3b.** Probability distribution for the frequency of conversion to AD, among the subpopulation having Olivia's, Ariel's, or Bianca's predictors. The red vertical line is the value of the probability p(cAD = Y | predictors).

shows the full-population distribution as comparison (dashed grey); with 95% probability Curtis's value is between 2.8 and 5.3, with a median of 3.8. And what is the frequency of conversion to Alzheimer's Disease among the subpopulation having Olivia's, Ariel's, or Bianca's predictors? The answer is given in the histogram of fig. 3b: with 95% probability, the fraction of this subpopulation that eventually converts to Alzheimer's Disease is between 0.19 and 0.43; this uncertainty range is due to the limited size of the learning dataset. The probability $p(CAD = Y \mid predictors)$ is equal to the average of such a distribution (e.g. Bernardo and Smith, 2000, §§ 4.2–4.3), provided the patient and dataset can be considered as coming from the same population.

Assessment of relevant subpopulation and auxiliary information 3.2

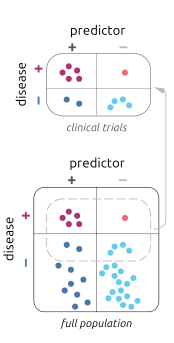
Rationale

As already mentioned, and as will be argued more concretely the next section, the clinician needs a probability in order to choose a treatment or other course of action about the current patient. This probability is computed by generalizing associations between predictors and predictand hidden in a dataset of similar patients, as discussed in § 2. The way this generalization is made, however, can differ from patient to patient in two respects:

- Only some particular directed associations can be generalized to the current patient, whereas others would be inappropriate to generalize. In some cases, for example when the learning dataset is artificially assembled with balancing or stratification methods, some associations cannot be generalized to any patients at all.
- There can be additional information available for the current patient, for instance some clinical predictors not included in the learning dataset, or other "softer" information such as family history or geographic background.

There is no sharp separation between these two items. The presence of additional information often automatically implies that some associations cannot be generalized from the learning dataset to the current patient.

Let us explain with a familiar example why particular associations cannot be generalized. Most medicine students learn about the *base-rate fallacy* (Bar-Hillel, 1980; Jenny et al., 2018; Sprenger and Weinberger, 2021; Matthews, 1996). Consider a large set of clinical trials, illustrated in the upper table on the side, where each dot represents, say, 10 000 patients. In this sample dataset it is found that, among patients having a particular value "+" of some predictors (left column), 71.4% of them (or 5/7, upper square) eventually developed a disease. The fallacy lies in judging that a new real patient from the full population, who has predictor value "+", also has a 71.4% probability of developing that disease. In fact, *this probability will in general be different*. In our example, it is 33.3% (5/15), as can be seen in the lower table illustrating the full population. This difference would be noticed as soon as the inappropriate probability was used to make prognoses in the full population. A similar situation happens for the predictor value "-".



There is a discrepancy in the conditional frequencies of predictand given predictors, between the sample dataset and the full population, because the proportion of positive vs negative disease cases in the latter has some value, 16.7%/83.3% in our example, whereas the samples for the trials (dashed line in the lower table) were hand-chosen so as to have a 50%/50% proportion. This sampling procedure is called "class balancing" in machine learning (Provost, 2000; Drummond and Holte, 2005; Weiss and Provost, 2003). More generally this discrepancy can appear whenever a population and a sample dataset from it do not have the same frequency distribution for the predictand. In this case, we cannot rely on the probabilities of "predictand given predictors" obtained from the sample dataset, which we symbolically write as

$$p(predictand | predictors, dataset)$$
 (1)

A little counting in the side figure reveals, however, that other frequencies may be relied upon. Consider the full population. Among all patients who developed the disease, 83.3% of them (or 5/6, upper row) had the predictor value "+", while among those who did not develop the disease, 33.3% (or 1/3, lower row) had the predictor value "-". And these frequencies are the same in the sample dataset. These frequencies from the clinical trials can therefore be used to make a prognosis using Bayes's theorem. For brevity, denote the predictors by X, the predictand by Y, the dataset or trials by D, and the full population by F. Bayes's theorem yields

$$p(Y \mid X, D, F) = \frac{p(X \mid Y, D) \cdot p(Y \mid F)}{\sum_{Y} p(X \mid Y, D) \cdot p(Y \mid F)}$$
(2)

Table 3. superiority of the "predictors | predictand" approach

We split our learning dataset in two subsets:

- One with 361 datapoints and a ratio of 29.9%/70.1% of conversions to Alzheimer's Disease vs stable Mild Cognitive Impairment. This is used as learning set.
- One with 343 datapoints and a ratio of 63.3%/36.7% of conversions to Alzheimer's Disease vs stable Mild Cognitive Impairment. This is used as a fictive full population.

This partition was made with no systematic sampling of any variates except the conversion variate. We then make a prognosis for each of the 343 "new" patients, through four separate approaches: (a) using the probabilities p(predictand | predictors, dataset), as typical of machine-learning algorithms; (b) using p(predictors | predictand, dataset) together with the base rate, as explained above; (c) tossing a coin; (d) always prognosing "conversion to Alzheimer's Disease", which guarantees 63.3% correct prognoses owing to the base rate. Finally, the accuracies (number of prognoses giving more than 50% probability to the correct course) of these four approaches are calculated. Here are the results from lowest to highest:

The "predictand | predictors" approach leads to worse results than a coin toss because of its underlying base-rate fallacy. The "predictors | predictand" approach leads to better results than simply always prognosing the most common base-rate outcome; this shows that the dataset can still provide useful statistical information despite its mismatched base rate. Inference algorithms that only yield "predictand | predictors" outputs, unlike the Ledley-Jaynes machine, are incapable of extracting this useful information.

In our example we find

$$p(Y = + | X = +, D, F) = \frac{p(X = + | Y = +, D) \cdot p(Y = + | F)}{p(X = + | Y = +, D) \cdot p(Y = + | F)p(X = + | Y = -, D) \cdot p(Y = - | F)}$$

$$\approx \frac{0.833 \cdot 0.167}{0.833 \cdot 0.167 + 0.333 \cdot 0.833} = 0.33$$
(3)

which is indeed the correct full-population frequency.

If the samples of the clinical trials had been chosen with the same frequencies as the full population (no "class balancing"), then the probability p(predictand | predictors, dataset) from the dataset would be the appropriate one to use. But the probabilities p(predictors | predictand, dataset) together with Bayes's theorem as in eq. (2) would also lead to exactly the same probability. We thus see that using the probabilities

$$p(\texttt{predictors} \mid \texttt{predictand}, \texttt{dataset})$$

from the dataset is preferable to using $p(predictand \mid predictors, dataset)$. The former yield the same results as the latter when use of the latter is appropriate, and allow us to apply corrections when use of latter is inappropriate. The superiority of using $p(predictors \mid predictand, dataset)$ probabilities is illustrated with a toy example in table 3.

The use of dataset probabilities different from p(predictand | predictors, dataset) can be necessary even when the dataset has statistics identical with the population it is sampled from. Typical cases are the prognosis of a patient that comes from a peculiar subpopulation or even from a different population (Lindley and Novick 1981; Quintana et al. 2017; Sox et al. 2013, ch. 4; Hunink et al. 2014, ch. 5). The first case happens for instance when the clinician has additional information not included among the predictor variates, such as the result of an additional clinical test, or family history. The second case happens for instance when the patient comes from a different geographical region. There is of course no sharp distinction between these two cases.

What is important is that in either case, it can still be possible to use statistical information from the sample dataset to make prognoses. It is sufficient that some *conditional* statistics may be applicable to the specific patient. For a patient coming from a different region, for example, it may be judged that the conditional probabilities p(predictand | predictors, dataset) still apply. In other words, the patient may still be considered of a member of the subpopulation having those specific predictor values. Using more technical language we say that a new patient can be considered *exchangeable* with the patients constituting the dataset, but only conditional on particular variates. See Lindley (2014, especially around §§ 7.3, 8.6; 1981) for a clear and logically impeccable presentation not obscured by technical language (more technical references are de Finetti 1930, 1937; Dawid 2013; Bernardo and Smith 2000, §§ 4.2–4.3; see also Malinas and Bigelow 2016, Sprenger and Weinberger 2021 about confounding and Simpson's paradox, to which this topic is tightly related).

This topic is complex and of extreme importance for inference, but its detailed study is not the goal of the present work. Our main point here is that population variability and auxiliary clinical information are important factors that differentiate patients, and a personalized approach ought to take them into account. The method here presented does this naturally, allowing a great flexibility in selecting which statistical features of the sample dataset should be used for each new patient, and the integration of auxiliary clinical information in the form of a prior probability. As discussed in § 3.1, the Ledley-Jaynes machine allows us to quickly calculate conditional probabilities $p(Y \mid X, \text{dataset})$ for any desired variate subsets Y and X required by the patient's relevant population.

Application to the example study

In our example of § 1.0, all statistics of the dataset are considered relevant for Olivia, Bianca, and Curtis. For these patients the clinician can therefore use Bayes's theorem with the likelihoods of table 2 and the dataset conversion rate of 0.463 – or equivalently directly the probabilities $p(CAD = Y \mid predictors, dataset)$ provided in the same table.

For Ariel, however, the clinician judges that a different base rate or prior probability of conversion should be used, equal to 65%, because of her different geographical origin and family history. In her case the clinician uses Bayes's theorem with the likelihoods of table 2 and the prior probability of 0.65.

The final probabilities of conversion to Alzheimer's Disease for our four patients are reported in table 4. Note how the final probability for Ariel is higher than that for Olivia and Bianca, even if the predictor data are the same for these three patients.

	Olivia	Ariel	Bianca	Curtis
initial probability $p(CAD = Y \mid auxiliaryinfo)$	0.463	0.65	0.463	0.463
$final\ probability\ p(\texttt{cAD} = \texttt{Y} \mid \texttt{predictors}, \texttt{dataset}, \texttt{auxiliaryinfo})$	0.302	0.47	0.302	0.703

Table 4. Final probabilities of conversion computed from dataset and auxiliary information

3.3 Assessments of treatments and benefits; final decision

Rationale

A crucial point in clinical decision-making is this: the clinician needs to assess, not the presence (present or future) of a disease, but the *risk* of its presence. Is there a difference between these two problems? and why is the difference important?

In clinical practice, we can rarely diagnose or prognose a medical condition with full certainty. Perfect classification is therefore impossible. But also a "most probable" classification, which may be enough in other contexts, is inadequate in clinical ones. The problem is that the clinician has to decide among different courses of action, such as different treatments, more tests, and so on, and the optimal one depends on *how probable* the medical condition is, not just on whether it is more probable than not.

Two examples illustrate this point. Suppose there is a dangerous treatment that extends the patient's lifetime by one year if the disease is on its course, but shortens the patient's lifetime by five years if the disease is not present. Also suppose that some algorithm tells the clinician whether the disease's presence is "more probable than not", given some predictor values; in which case the clinician administers the dangerous treatment. It turns out that 60 out of 100 treated patients having these predictor values eventually develop the disease, so "more probable than not" is correct. However, the clinician has added $1 \times 60 = 60$ years but subtracted $5 \times 40 = 240$ years from the combined lifespans of the treated patients! The conclusion is that the treatment cannot be prescribed just because the disease is "more probably present than not". As an opposite example, suppose that a less dangerous treatment extends the patient's lifespan by five years if the disease is on its course, but shortens it by one month if the disease is not present. In this case, it may be advisable to undergo the treatment even if the disease is less probably present than not. If the clinician administer the treatment to 100 similar patients, and 20 of them develop the disease, then the clinician has added $5 \times 20 = 100$ and subtracted $\frac{1}{12} \times 60 = 5$ years to their combined lifespans.

In both examples, it is clearly important to assess the *probability* – having precise connections with the population frequency – that the patient will develop the disease. ¹³ The Ledley-Jaynes machine, as explained in the previous sections, gives us such probabilities.

But the choice between treatments depends not only on the probability of the medical condition. Here is where differences between patients vary and matter the most. Consider again the second example above, about the less dangerous treatment. Let us add that the treatment would extend the lifespan by five years, but would also somewhat worsen the quality of life of the patient and of the patient's family. Suppose our patient is quite old and tired, has had a happy life, and is now looking with a peaceful mind towards death as a natural part of life. Such a patient may prefer to forego the bother of the treatment and the additional five years, even if the probability for the disease is quite high.

¹³ In the first example, the treatment should only be administered if the probability is higher than 83.3%; in the second, it can be administered if the probability is at least 1.6%.

The benefits of the different treatments, and the probability thresholds at which one treatment becomes preferable to another, must therefore be judged and quantified primarily by the patient. Utility theory and maximization of expected utility allow clinician and patient to make such judgements and decisions in a coherent way (Sox et al. 2013; Hunink et al. 2014; see also the clear and charming exposition by Lindley 1988, and O'Hagan et al. 2006).

We summarize the main, patient-dependent procedure for decision-making, and show how our computations so far fit perfectly with it.

The clinician first assesses and lists the mutually exclusive courses of actions available for the specific patient. These could be treatments, more tests, doing nothing, and so on. Often there are *sequences* of decision available, but the utility framework can be applied to them as well (see references above and Raiffa, 1970). The list of courses of action is already patient-dependent: some alternatives may not be suitable (say, owing to allergies), some may be economically too costly, and so on.

Each course of action will have different consequences, which additionally depend on the patient's unknown clinical condition of interest. A treatment may have some consequences if the patient has or will develop the disease, and different consequences otherwise. The patient quantifies, with the clinician's guidance, the benefits and costs – technically called "utilities" – of such possible consequences. The quantification of utilities is not within the scope of the present work. The references cited above offer several guidelines and rules for numerically translating factors such as quality of life and expected lifespan into utilities.

The courses of actions, uncertain clinical conditions, and the quantified utilities U of their consequences can be organized into a table of this form:

	clinical condition a	clinical condition b	
action α	$U_{lpha a}$	$U_{\alpha b}$	
action β	$U_{eta a}$	$U_{eta b}$	

which can be compactly represented by a so-called *utility matrix* (U_{ij}) , the row index i enumerating the actions, and the column index j the clinical conditions. Note that the number of possible treatments and clinical conditions do not need to be equal; generally, they are not.

The expected utility \bar{U}_i of an action i is calculated as the expectation of its utilities U_{ia}, U_{ib}, \ldots with respect to the probabilities $p(a), p(b), \ldots$ of the clinical conditions a, b, \ldots :

$$\bar{U}_i := U_{ia} \, \mathrm{p}(a) + U_{ib} \, \mathrm{p}(b) + \cdots \tag{4}$$

Note that this corresponds to a matrix multiplication between the matrix of utilities and the vector of probabilities.

Finally, the recommended action is the one having *maximal expected utility*.

Application to the example study

At present, there are no cures for Alzheimer's Disease, although some recent pharmacological agents are shown to extend the time before a patient is cognitively severely impaired * ref. But for the sake of our

case study let us imagine that there, in the near future, are three mutually exclusive treatment options for prevention or retardation of the disease; call them β , γ , δ . And denote the simple option of "no treatment" by α . The clinical conditions to be considered are just two: the patient will convert to Alzheimer's Disease (CAD = Y), or will remain with stable Mild Cognitive Impairment (CAD = N).

We have therefore $4 \times 2 = 8$ possible consequences of the four treatments depending on the two clinical conditions. Our four patients and clinician quantify the utilities, arriving at the utility matrices shown in table 5, top. Olivia, Ariel, and Curtis quantify the benefits of the treatments in exactly the same way, but Bianca's quantification differs slightly, because of the interaction of the treatments with several allergies and additional clinical conditions, as explained in § 1.0.

Utility matrices						
	Olivia	Ariel	Bianca	Curtis		
$\begin{array}{c} \text{treatment } \alpha \\ \text{treatment } \beta \\ \text{treatment } \gamma \\ \text{treatment } \delta \end{array}$	$\begin{bmatrix} 10 & 0 \\ 9 & 3 \\ 8 & 5 \\ 0 & 10 \end{bmatrix}$	$\begin{bmatrix} 10 & 0 \\ 9 & 3 \\ 8 & 5 \\ 0 & 10 \end{bmatrix}$	$\begin{bmatrix} 10 & 0 \\ 8 & 3 \\ 7 & 5 \\ 0 & 10 \end{bmatrix}$	$\begin{bmatrix} 10 & 0 \\ 9 & 3 \\ 8 & 5 \\ 0 & 10 \end{bmatrix}$		

Expected utilities and optimal treatments					
	Olivia	Ariel	Bianca	Curtis	
treatment α treatment β	6.98 7.19	$5.27 \\ 6.16$	6.98 6.49	$\frac{2.97}{4.78}$	
treatment γ	7.09	6.58	6.40	5.89	
treatment δ	3.02	4.73	3.02	7.03	
<u>optimal</u>	β	γ	α	δ	

Table 5. Utility matrices, expected utilities, and optimal treatements for our four patients

The probabilities for the two medical conditions are those found in the previous subsection, table 4. For brevity, we denote just by p(cAD) the probability of conversion given a patient's predictor values, and by $p(\texttt{sMCI}) \equiv 1 - p(\texttt{cAD})$ the complementary probability of stable Mild Cognitive Impairment, given the same predictor values. The expected utilities of each treatment for each patient can then be easily computed. For example, for Olivia the expected utility of treatment β is

$$\bar{U}_{\beta} = 9 \cdot (1 - 0.463) + 3 \cdot 0.463 = 7.19$$
 (5)

The results for all patients are reported in table 5, bottom, with the maximal expected utilities in **boldface**.

3.4 Additional information provided by the Ledley-Jaynes machine

As already said, the output of the Ledley-Jaynes machine is about the full population of future patients, with all its statistics. This output can therefore be used for additional purposes such as resource planning, imputation of missing data, sensitivity checks, and the investigation of the predictors' importance in the prognosis.

Resource planning

Let us ask the following question: if the learning dataset were representative of the full population, then how often, in the long run, would a clinician prognose a conversion to Alzheimer's Disease with probability between 0%-2%, or 2%-4%, and so on with 50 bins up to 98%-100%?

The Ledley-Jaynes machine can answer this question probabilistically; the answer is plotted in fig. 4. Note that the calculation assumes that the Ledley-Jaynes machine will not be regularly updated with new patients' data (the calculation could also be made with the opposite assumption). The light-blue bands are 95% uncertainty coverage intervals ¹⁴; this uncertainty comes from the fact that we are not certain about the full-population frequencies. We see that it is very improbable that many patients will be prognosed with probabilities around 30%, smaller than 5%, or larger than 75%. The full population is likely to be grouped into four or five "conversion-probability clusters", as evident from the peaks.

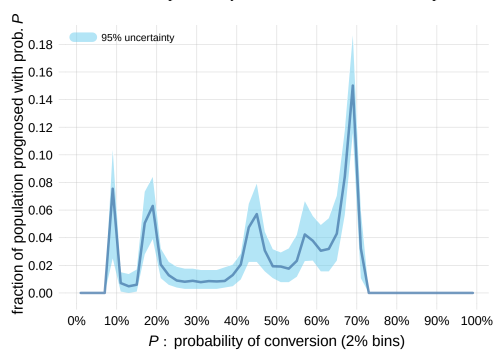


Figure 4. Possible distribution of prognostic probabilities of conversion in the full patient population

This kind of information allows us to make other forecasts of various kinds. For instance, it would be useful to forecast in which proportions the treatments α , β , γ , δ (see § 3.3) will be prescribed, assuming that the full population has on average the utility matrix of Olivia, table 5. We find these coverage intervals with a 90% probability that the true future proportion will be within each:

$$\alpha: 21\%-28\%$$
 $\beta: 2\%-6\%$ $\gamma: 31\%-42\%$ $\delta: 31\%-40\%$ (6)

Again, despite the obvious uncertainties, we can be quite sure that treatments γ , δ will be prescribed more often than α , and that β will be only prescribed in 5% of cases. This kind of semi-quantitative forecast can be very useful for resource planning.

¹⁴ for terminology see JCGM (2008, C.2.30). A p% coverage interval or credible interval is an interval containing the true value with p% probability. Note that it is different from a "confidence interval", which cannot be interpreted in such a simple way (Pratt, 1961, pp. 165–166; Jaynes, 1976; MacKay, 2005, § 37.3)

Imputation of missing data

As mentioned in § 3.1, the Ledley-Jaynes machine treats all variates of the learning dataset on the same footing. This is why it can output probabilities "predictand | predictors", "predictors | predictand", and other combinations with equal ease, exploiting them to correct subpopulation mismatches as illustrated in § 3.2.

This feature allows us to impute missing data for one or more patients, giving a probability distribution of what those data could be. Such imputation can be done at prognostic time, for sensitivity checks; we discuss this more in detail in the next section. The imputation can also be done a posteriori, possibly years later, when the actual predictand value becomes known. This can be useful for many purposes, for instance, for the comparison of biological hypotheses.

Sensitivity checks

The imputation of missing data at prognostic time is useful for various kinds of sensitivity analysis.

Let us consider for instance the case of Curtis, whose value of Hippocampal Volume is missing (table 2). His clinician thus wonders if the acquisition of this value could lead to a different treatment. The Ledley-Jaynes machine can give a probabilistic answer to this question. We already saw in § 3.1, fig. 3a, that the machine can tell the clinician what are the probable values of this predictor for Curtis.

The clinician can also use the Ledley-Jaynes machine to find the probability distribution of Curtis's expected utilities (table 5) *if the Hippocampal Volume had been known*. The result is that the expected utilities for the four treatments in Curtis's case must be in the following coverage intervals with 90% probability:

$$\alpha$$
: 2.97%–2.98% β : 4.78%–4.79% γ : 5.89%–5.90% δ : 7.02%–7.03% (7)

(the four corresponding probability histograms, if plotted jointly, would look like distinct vertical lines). It is clear that knowledge of the Hippocampal Volume would be overly unlikely to change Curtis's optimal treatment from δ . The clinician therefore proceeds without this predictor.

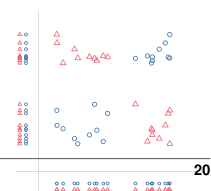
Predictor importance

Luca] I think the initial presentation of the problem, here below, still slightly misses the target. The point is that the vague and ill-posed question "which predictor is most important?" hides a decision problem: "if I had to drop one feature for all patients, which one should I drop last?" or "if I had to use only one feature for all patients, which one should I prioritize?". This formulation uniquely determines the relevant metric. I'll rewrite this later.

The question about Curtis in the previous subsection can be generalized to a whole population. How important, in general, is each predictor in prognosing the conversion to Alzheimer's Disease? Predictors that are too invasive or too expensive to obtain and do not make a real difference in the prognosis could be dropped altogether.

To answer this general question, which is too vague (ill-posed), we must first specify which other predictors the predictor of interest is used with, and what we mean by "important".

The schematic picture on the side illustrates the necessity of specifying a predictor's context. Individuals in this population can be $\stackrel{\aleph}{\sim}$ either blue circles \circ or red triangles \triangle and have two predictors X_1, X_2 .



Predictor X_1 , if used by itself, is worthless in distinguishing the two subpopulations, because these have identical marginal distributions (depicted beside the grey lines). If used in conjunction with X_2 , however, predictor X_1 allows us to identify an individual's subpopulation with full certainty. It is therefore an essential predictor in this case:

dropping it would lead to a complete loss of predictive power. An analogous discussion holds for X_2 in the present case. The converse can also happen (not illustrated): a predictor might be "good" if used by itself, and yet it might be discarded without any loss if used in combination with others.

A predictor's "importance" or "prognostic power" is undefined until we specify a relevant metric, for predictors can be ranked differently by different metrics. From our discussion so far it should be clear that in clinical decision-making the canonical metric is the *utility* which a predictor's presence or absence leads to. This point was partially illustrated with Curtis's example in the previous subsection. What if we want to make an assessment, not for a single patient, but for the full population? It can be proved, again from decision-theoretic principles, that the population average of all utility matrices should be used in this case (cf. Dyrland et al., 2022a, § 4.1). This seems a quantity very difficult to assess; but it can also be proved that even just a semi-quantitative assessment leads to better results than using some other "general-purpose" metric (Dyrland et al., 2022a, § 4.2).

The Ledley-Jaynes machine allows us to compute the expected value of virtually any prognostic-importance metric of any subset of predictors available in the dataset. Two facts of paramount advantage are that (a) the prognostic power of a set of predictors found with the Ledley-Jaynes machine is the maximal one obtainable by any inference algorithm, or in other words it is an intrinsic property of that set of predictors; (b) the Ledley-Jaynes machine achieves this maximal power. Thus, if the Ledley-Jaynes machine says that the accuracy obtainable with a given set of predictors is 70%, then we know that no other inference algorithm of any kind can reach a higher accuracy than 70%; inference algorithms that reach lower accuracy can in principle be improved upon. The Ledley-Jaynes machine, by construction, will reach this accuracy.

We illustrate this kind of computation in our example case, using: (a) two metrics, the accuracy and the mutual information (Shannon, 1948; Cover and Thomas, 2006) between a set of predictors and the event of conversion to Alzheimer's Disease; and (b) 27 different sets of predictors:

- every predictor, used individually (12 sets);
- all cognitive-test predictors used together, jointly with demographic information (Age and Sex);
- APOE4, Hippocampal Volume, and demographic information, used together;
- all predictors together minus one, excluding each single predictor in turn (12 sets);
- all predictors jointly.

Use of the accuracy assumes that the population of patients has only two available treatments having average utility matrix $\begin{bmatrix} 1 & 0 \\ 0 & 1 \end{bmatrix}$. Mutual information is a model-free measure of the relation between two sets of variates, with diverse operational interpretations (MacKay, 2005; Woodward, 1964; Minka, 2003; Good and Toulmin, 1968; Kelly, 1956; Kullback, 1978) and international standards (ISO, 2008). A set of predictors and a binary variate (such as our conversion to Alzheimer's Disease) have a mutual information of 1 Sh if and only if there is a non-constant deterministic function from the former to the latter.

Our specific questions are the following: "What is the expected value of the accuracy for the next new patient, if we use the given set of predictors?" and "What is the mutual information between the given set of predictors and the predictand, given the presently available data?".

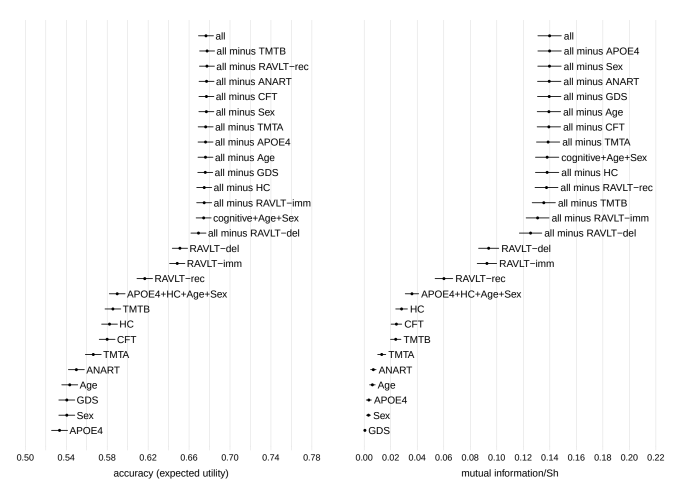


Figure 5. Expected accuracy (for the next new patient) and mutual information of several sets of predictors for the prognosis of conversion to Alzheimer's Disease. Both graphs have been vertically ordered according to increasing values; the two rankings agree within the respective uncertainties. The all predictor set is mathematically guaranteed to be optimal according to both metrics and has therefore been ranked first. Bars show the uncertainty interval (\pm two standard deviations).

The answers to these questions are reported in fig. 5, ordered from bottom to top according to increasing metric. The ordering of mutual information and accuracy agree within the uncertainty of the numerical computation (Monte Carlo integration). The latter is reported as coverage intervals of \pm two standard deviations.

The figure reveals several interesting findings, *valid within the population selected for the dataset*, which can be compared with the analysis in Rye et al. (2022, see especially Fig. 3 and Table 3):

- The set of 12 predictors considered in the present work and in Rye et al. (2022) can, at most, lead to a prognostic accuracy of around 68%, for any inference algorithm. This fact agrees with the (completely independent) findings in Rye et al. (2022), where a maximal accuracy of 68% was found using an ensemble model; the present analysis also shows that that model managed to achieve the maximal accuracy possible with these predictors (but see § 4 for limitations of that model.)
- APOE4, GDS, Age, Sex, and to some degree ANARTare poor predictors (within this population) when used alone and when used in combination with all other predictors. The latter point is evident from the fact that the mutual information of the combined predictors barely decreases if any one of these four predictors is omitted.

- The combined cognitive and demographic variates are better predictors than the combined hippocampal, APOE4, and demographic variates.
- RAVLT-imm, RAVLT-del, and to a lesser degree are good predictors, both when used alone and when used in combination with all other predictors. Hippocampal Volume is a poorer predictor than any of the RAVLT when used alone, and likely also when used in combination with all others (contrast this with Rye et al., 2022).

The Ledley-Jaynes machine shows that the omission of any one of the 12 predictors, except RAVLT-del and possibly RAVLT-imm, does not lead to an appreciable decrease in accuracy (relative decrease of 0.3% or less) or in mutual information (relative decrease of less than 3%). This puts the importance analysis of Rye et al. (2022) into perspective. The exact quantification of these subtle differences is computationally quite expensive, and we did not carry it out further.

3.5 Summary of the case study

Summary in table 6.

4 DISCUSSION

[Luca] There are a couple different ways to introduce the discussion. I'll write some variants as soon as possible

The differences among patients which are relevant to clinical diagnosis/prognosis and decision-making can be approximately divided into three categories:

- 1. differences in the availability and values of a core set of clinical predictors, for which we have population-wide statistical information;
- 2. differences in the availability and values of auxiliary and usually "softer" predictors, such as geographical or family background; or more generally of predictors not belonging to the core set;
- 3. differences in the availability and benefit of clinical courses of actions, such as treatments and further tests.

These differences are intimately connected with, and partially stem from, several hallmarks of a clinician's tasks:

First, there is often an irreducible element of uncertainty in inferring a medical or biological condition from a set of predictors. In mathematical terms, there is no function from predictors to predictands or refer to a figure in the previous sections. This is especially often the case for predictors that are more readily available and less invasive and, therefore, more desirable. The irreducible uncertainty originates from the natural variability within a population. Yet this variability can be harnessed, and the way to harness it can crucially vary from patient to patient. We illustrated this with an example in the inset of § 3.2, p. 14, which showed that any relation from predictors to predictands lead to poor prognoses, whereas taking into account the variability in the predictors given the predictand improved our prognoses.

Second, owing to the irreducible prognostic uncertainty, the clinician's ultimate task is not one of classification or regression, but one of *decision-making under risk*. This is clear if we consider that a clinician may have *three or more* courses of action to choose from, even if the unknown medical condition only has *two* possible "class labels". A pure class label is, therefore, of no help to the clinician, who

	Olivia	Ariel	Bianca	Curtis
Predictor values				
Age Sex HV/10 ⁻³ APOE4 ANART CFT GDS RAVLT-imm RAVLT-del RAVLT-rec TMTA TMTB	75.4 F 4.26 N 18 21 3 36 5 10 21 114	75.4 F 4.26 N 18 21 3 36 5 10 21	75.4 F 4.26 N 18 21 3 36 5 10 21	63.8 M [missing] Y 15 14 2 20 0 3 36 126
Additional information, final probability auxiliary info applicable dataset statistics prior probability of conversion	none all 0.463	family history, base rate predictor predictand 0.65	none all 0.463	none all 0.463
Available actions and utilities $ \begin{array}{c} \text{sMCI cAD} \\ \text{treatment } \alpha \\ \text{treatment } \gamma \\ \text{treatment } \delta \end{array} $	$\begin{bmatrix} 10 & 0 \\ 9 & 3 \\ 8 & 5 \\ 0 & 10 \end{bmatrix}$	$ \begin{bmatrix} 10 & 0 \\ 9 & 3 \\ 8 & 5 \\ 0 & 10 \end{bmatrix} $	$ \begin{bmatrix} 10 & 0 \\ 8 & 3 \\ 7 & 5 \\ 0 & 10 \end{bmatrix} $	$ \begin{bmatrix} 10 & 0 \\ 9 & 3 \\ 8 & 5 \\ 0 & 10 \end{bmatrix} $
Outputs of Ledley-Jaynes machine				
$\begin{array}{l} p({\rm AD} \mid {\rm predictors}) \\ p({\rm predictors} \mid {\rm cAD})/10^{-12} \\ p({\rm predictors} \mid {\rm sMCI})/10^{-12} \end{array}$	0.302 8.97 18.6	0.302 8.97 18.6	0.302 8.97 18.6	0.703 1.14 0.343
final probability of conversion	0.302	0.47	0.302	0.703
exp. utility treatment α exp. utility treatment β exp. utility treatment γ exp. utility treatment δ	6.98 7.19 7.09 3.02	5.27 6.16 6.58 4.73	6.98 6.49 6.40 3.02	2.97 4.78 5.89 7.03
Optimal treatment	\boldsymbol{eta}	γ	α	δ

Table 6. Summary. The basic data that distinguish Ariel, Bianca, Curtis from Olivia are in red.

instead needs to know the uncertainty or probability of the label, in order to evaluate risks and benefits, and from these, make a final decision. The choice of a course of action is moreover the moment where patient differences can be very dramatic.

to be finished

Possible additional point of view Let's assume that current technology allows for a particular maximal rate of extraction of information per unit time, and if we assume that different inference algorithms reach this maximum rate (or do not reach it by a common amount). Information and communication theory

(MacKay, 2005, I–II) then suggests that if one inference algorithm is, say, 100 times faster than a Ledley-Jaynes machine, its output has also 100 times less information than a Ledley-Jaynes machine's. Do we need this extra information? The example case illustrated in the present work shows that part of this extra information is necessary for personalized medicine; and the remaining amount can be very useful or, at times, necessary for sensitivity analyses or resource planning.

4.1 Counters to critiques

Any inference or decision-making algorithm aspiring to take into account patient differences must perforce have some open "input slots" for such differences. We saw that the Ledley-Jaynes machine requires inputs about a patient's specific predictors, relevant statistical relations and auxiliary data, and treatment utilities.

Some researchers may wonder: can such additional input be avoided? They fear that errors could sneak in through these extra inputs.

This question is answered by a mathematical theorem at the very core of decision theory ¹⁵, which is too seldom emphasized: Any decision we make, either (A) comes explicitly or implicitly through some set of utilities and maximization of their expectations, or (B) is logically inconsistent. There is no third alternative. Thus the choice is not between using utilities or not using utilities, but between choosing them explicitly or letting them be chosen in a way we don't know. If you use a decision-making algorithm that does not ask you for utilities, then the algorithm is internally supplying utilities beyond your control (and probably divorced from your specific problem), or, worse, is committing logical inconsistencies.

The first advantage of explicitly operating through utilities, probabilities, decision theory, is that we are, at the very least, sure of not acting in a self-contradictory way. The second advantage is that the utilities used to arrive at a decision lie openly in front of us. We can analyse and change them if we find them inappropriate to a specific problem. If they are hidden, it's more difficult to analyse which are inappropriate and how they should be changed.

Subtly hidden disastrous consequences of not following normative decision theory: An algorithm can lead to saving 85 000 patients out of 100 000 and be deemed a success. But if the ideal algorithm had been used, 95 000 patients would actually have been saved. What shall we say to the families of the 10 000 patients who could have been saved but weren't?

4.2 Range of application of Ledley-Jaynes machines

The range of application of the Ledley-Jaynes machine used in the present study has two kinds of bound: computational and theoretical.

As mentioned in § 3.1 and explained in appendix A.1, the fact that the Ledley-Jaynes machine extracts all available information from the dataset also makes it computationally expensive. It is at present impossible to use with high-dimensional predictors (if our dataset had included a predictor such as a $128 \times 128 \times 128$ grayscale MRI image, the learning stage would have taken around 100 years). Approximate but much faster algorithms such as neural networks and random forests are thus, at present, still the only options with such predictors. There is, however, the interesting possibility of combining such fast algorithms together with a Ledley-Jaynes machine, as a post-processor of their raw output. This allows us to extract useful information

¹⁵ Savage (1972); Luce and Raiffa (1957); Raiffa and Schlaifer (2000); Atkinson et al. (1964); Ferguson (1967); Lindley (1988); Kreps (1988); Bernardo and Smith (2000); Pratt et al. (1996); Lindley (2014); Pettigrew (2019)

usually hidden in such output at a low computational cost (Dyrland et al., 2022b). Such information can be then used for clinical decision-making as illustrated in the present work.

As explained in § 3.2, the essentially sole assumption underlying the Ledley-Jaynes machine's inference and its practical use with new patients, is that the latter can be assumed to come, at least in some respects, from the same population as the learning dataset (in technical jargon, partial or conditional exchangeability applies). This precludes using the Ledley-Jaynes machine to forecast how the statistics of the full population could change in the future. However, the machine can be used for time-dependent (transversal correct?) inferences within a stable population, such as forecasts of the future time of disease onset, expected lifelength, and similar. For example, if data about the time of conversion to Alzheimer's Disease were available in the dataset, the Ledley-Jaynes machine could forecast not only whether, but also when the conversion could take place (cf. e.g. De la Cruz-Mesía et al., 2007).

A APPENDICES

A.1 Mathematical details about the Ledley-Jaynes machine

As discussed in § 3.1, the Ledley-Jaynes machine explores the space of possible distributions of frequencies of all 13 variates listed in table 2, for the full population of patients from which the dataset originates. In the present study, it does so by using a total of 1535 independent parameters to represent the distributions, with roughly 190 parameters for each continuous or integer variate. As a crude intuition, it is as if we divided the range of each variate into 190 bins, and considered all possible frequency histograms over these. The actual parametrization is smarter, using parameters to represent less and less smooth traits of the distribution. We indeed expect the distribution for a full population to have some degree of smoothness, owing to physical and biological reasons. Actually, the number of parameters used is in principle infinite, because the machine gives a warning if the data indicates that more parameters are needed. In the present study, the data indicate, on the contrary, that fewer than 250 parameters would be enough. More details on the mathematical representation can be found in Dunson and Bhattacharya (2011); see also Rossi (2014); Rasmussen (1999).

★ Should I add a more precise description of the mathematical representation?

There is a fundamental difference in how the Ledley-Jaynes machine and most popular machine-learning algorithms (including neural networks, random forests, support-vector machines, excluding gaussian processes) work. The latter do, at bottom, an optimization, looking for the minimum of some error function. The Ledley-Jaynes machine does a full *space exploration* and *averaging*, as explained in § 3.1. Inference and generalization in fact essentially rely on averaging operations in problems such as the present one (de Finetti 1930, 1937; Dawid 2013; Bernardo and Smith 2000, §§ 4.2–4.3; see also Self and Cheeseman 1987). The optimization done by most machine-learning algorithms is an approximate form of averaging – assuming or hoping that most of the mass to be averaged is around the extremum (MacKay, 1992a; Murphy, 2012, ch. 16). But the underlying necessity of a proper averaging becomes manifest in many of the obligatory procedures that go together with training a machine-learning algorithm; cross-validation for instance (MacKay, 1992b). Add note: "ensembling" in ML is not this kind of averaging (Murphy, 2022, § 18.2)

This difference explains why the Ledley-Jaynes machine is computationally much more expensive than other algorithms, but also why its output is informationally so rich, and why it does not need any validation datasets, test datasets, other data splits, or cross-validation procedures (it can be proved that one of the internal computations of the machine is mathematically equivalent to doing k-fold cross-validations for *all possible* data splits and k; see e.g. Porta Mana 2019; Fong and Holmes 2020).

one more remark about extremum-search being equivalent to making a choice, but the utilities are not controlled by the patient and not flexible.

A.2 Computational details

The learning stage, with 13 variates and 704 datapoints, took less than 5 hours. The computation was done on a 16-core Intel Core i9-12900K. After that, the calculation of probabilities and expected utilities for any single patient is immediate. The mutual information and accuracy analysis of § 3.4 took roughly 1 h.

CONFLICT OF INTEREST STATEMENT

The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

AUTHOR CONTRIBUTIONS

The authors were too immersed in the development of the present work to keep a detailed record of who did what.

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SUPPLEMENTAL DATA

Supplementary Material should be uploaded separately on submission, if there are Supplementary Figures, please include the caption in the same file as the figure. LaTeX Supplementary Material templates can be found in the Frontiers LaTeX folder.

DATA AVAILABILITY STATEMENT

The datasets [GENERATED/ANALYZED] for this study can be found in the [NAME OF REPOSITORY] [LINK].

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