Response heterogeneity: Challenges for personalised medicine and big data approaches in psychiatry and chronic pain [version 1; referees: 1 approved, 1 approved with reservations]

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Abstract
Response rates to available treatments for psychological and chronic pain disorders are poor, and there is a considerable burden of suffering and disability for patients, who often cycle through several rounds of ineffective treatment. As individuals presenting to the clinic with symptoms of these disorders are likely to be heterogeneous, there is considerable interest in the possibility that different constellations of signs could be used to identify subgroups of patients that might preferentially benefit from particular kinds of treatment. To this end, there has been a recent focus on the application of machine learning methods to attempt to identify sets of predictor variables (demographic, genetic, etc.) that could be used to target individuals towards treatments that are more likely to work for them in the first instance. Importantly, the training of such models generally relies on datasets where groups of individual predictor variables are labelled with a binary outcome category – usually ‘responder’ or ‘non-responder’ (to a particular treatment). However, as previously highlighted in other areas of medicine, there is a basic statistical problem in classifying as ‘responding’ to a particular treatment on the basis of data from conventional randomized controlled trials. Specifically, insufficient information on the partition of variance components in individual symptom changes mean that it is inappropriate to consider data from the active treatment arm alone in this way. This may be particularly problematic in the case of psychiatric and chronic pain symptom data, where both within-subject variability and measurement error are likely to be high. Here, we outline some possible solutions to this problem in terms of dataset design and machine learning methodology, and conclude that it is important to carefully consider the kind of inferences that particular training data are able to afford, especially in arenas where the potential clinical benefit is so large.
Introduction

The proportion of patients who respond to available treatments for psychological and chronic pain disorders is often low. For example, in major depression, roughly 40% of individuals experience a ‘clinically significant’ response (decrease in symptom severity score above some minimum value) over the course of treatment (e.g. 1,2). Similarly, a recent meta-analysis of available pharmacotherapies for neuropathic pain found estimates of ‘number needed to treat’ (number of patients needed to be treated to prevent one additional adverse clinical outcome) for effective treatments ranged from 4–10, indicating poor response rates.3 For patients, this often means a lengthy process of cycling through different treatment options, in a sequence that may be significantly influenced by non-clinical concerns (e.g. relative drug cost, therapist availability, local health authority guidelines), and where there may be inadequate data on the safety and effectiveness of switching regimes (e.g. 4). For psychological conditions, this process can be particularly lengthy, given the significant period of time before common pharmacological treatments are expected to take effect (e.g. 4–6 weeks to conclude a particular drug treatment is ineffective). Together, this results in a substantial burden of suffering and disability to individuals with a diagnosis of these disorders, before (if) an effective treatment option can be found.

It is generally assumed that differential response to a particular treatment across individuals can be at least partially explained by patient heterogeneity within a certain diagnostic category – i.e. that individuals who present to the clinic with similar sets of symptoms may have different underlying pathologies. This seems particularly reasonable assumption in the case of both mental health disorders and chronic pain, as diagnosis is often made purely on the basis of self-reported symptom checklists, and our lack of knowledge into the aetiology of these conditions means we have little opportunity for differential diagnosis. Indeed, in the case of psychiatric disorders, such as depression, diagnosis can often be made on the basis of directly contradictory symptom reports (e.g. sleeping too much vs. sleeping too little), and there may be many different ways to meet diagnostic criteria (e.g. 227 possible symptom combinations for major depressive disorder, according to DSM-IV). Similarly, even patients with a diagnosis of a particular pain condition are likely to have distinct patterns of nervous system damage, involving multiple pathways (e.g. 5), and definitions of chronic pain itself can vary dramatically across research groups and clinical centres.

Even if we lack insight into pathological mechanisms, it seems likely that if we are able to use some kind of predictive method to direct individuals towards treatments that are likely to be more effective for them – then even a small increase in the resulting response rate could potentially have a large effect on disease burden for individual patients. There has therefore recently been great interest in doing just this for psychiatric data, via application of supervised learning methods to large datasets of individual clinical predictors and treatment response data (see 8 for an excellent recent review of potential clinical advantages and best methodological practice in this area).

The current gold standard approach is firstly to define a set of features and targets for various machine learning algorithms to train on. In this context, features are individual difference variables that may potentially relate to future treatment outcome (clinical, demographic, physiological, genetic, behavioural, etc. information). The target variable (that the algorithm must learn to predict) is usually a binary category label, such as ‘responder’ or ‘non-responder’ (whether or not an individual has exhibited symptom improvement above some threshold level, following a particular course of treatment). Various supervised learning algorithms can then be trained on this labelled dataset (ideally using a rigorous cross-validated approach), and assessed in terms of their predictive accuracy on independent 'unseen' (during model training) data. Finally, the best model can be brought forward to a randomised controlled trial framework, where treatment allocation by current clinical guidelines could be compared to algorithm-assisted treatment assignment.

This approach is highly attractive, as the potential clinical gains from even a small increase in likelihood of treatment response for a particular individual are large. However, across the field of medicine in general, attempts to make such clinical gains via a personalised medicine approach have not often fulfilled their initial promise – with relatively few reaching the clinic (e.g. 9). Here, we explore a basic statistical issue that may limit the effectiveness of this process – i.e. the reliability of distinguishing between treatment ‘responders’ and ‘non-responders’ in the first place. We further discuss the reasons why this problem may be particularly acute in the case of available data regarding psychiatric disorders and chronic pain conditions, and some potential solutions.

The problem of response heterogeneity

The problem of properly identifying response heterogeneity, or, more simply, reliably distinguishing between responders and non-responders to a particular treatment, on the basis of randomised controlled trial (RCT) data, has previously been highlighted across various fields of medicine.10-12 If not properly addressed, this constitutes an absolute limit on the effectiveness of predictive models at the level of input or training data, thereby limiting their future clinical usefulness.

The issue is best illustrated by considering the nature of data collected during RCTs, and the kind of inferences this process affords. The foundation of an RCT is that the mean effect of an intervention (e.g. active drug treatment) is derived by comparing what happened, on average, to the (randomly allocated) participants in the intervention group to what happened, on average, to participants in the control (e.g. placebo) arm. The random allocation of participants to the intervention vs control arms allows the control group to function as an illustration of what we might have expected to occur in the intervention group, had they not received the active treatment – in turn allowing us to draw conclusions about the overall (average) effects of the treatment itself.13 Crucially, we can only draw this inference by direct comparison to the control arm data.

This basis of an RCT means that we cannot identify responders and non-responders by considering individuals in the intervention group alone. In other words, we cannot legitimately label an individual who received a particular active treatment as a ‘responder’ (or not) because we do not know what would have happened to
that particular individual if they had been in the comparator (or placebo) arm. This kind of information is very hard to obtain at the individual (cf the group) level, as there is no good way to obtain a control observation. Formally, to properly infer whether a particular participant responded or didn’t respond to a particular treatment, we would require knowledge of what would have happened if a key event (treatment administration) both did and did not occur (a form of counterfactual reasoning), which is not possible in the real world.

A particularly acute issue for psychiatric and chronic pain datasets?
Variability of change (e.g. \( t_2 - t_1 \) symptom score) in the intervention arm is not a true estimate of variability in treatment response, because it includes components of within-subject variation and measurement error. Even if measurement error is small (i.e. we can precisely measure the outcome variable of interest), for many medical interventions, the outcome variable will depend on a complex interplay of biological factors (e.g. time of day, stress level, etc.), and so within-subject variability will be relatively high. This means that the reliability of within-subject measurements across time points can be somewhat poor, and large variation in changes between study time points may be evident – even where there is no true individual difference in treatment response.

Unfortunately, for psychiatric and chronic pain symptom data, both measurement error and within-subject variation are likely to be high. Measurement error may be higher than other areas of medicine, as the main tools used to assess clinical outcomes are patient or clinician-completed questionnaire measures, which are relatively low precision tools. Further, although self-reported symptom levels are considered the gold standard outcome measure for both psychiatric disorders and chronic pain conditions, reliability is limited by factors such as cognitive capacity and level of insight for patient-rated measures (e.g. 14), and by interviewer skill and inter-rater agreement for clinician-rated measures (e.g. 15–17). Finally, these classes of disorders represent episodic, chronically relapsing conditions, which will likely contribute to large within-subject variation, particularly at typical RCT follow-up timescales (often around 6 months–1 year; cf e.g. median duration of a depressive episode of ~20 weeks.). If the variation in outcome due to these sources is greater than that due to any true individual differences in treatment response, it will be very hard to detect the latter under a conventional RCT framework.

A further problem in predicting true response heterogeneity is susceptibility of symptom change data to regression to the mean and mathematical coupling artefacts. Regression to the mean refers to the phenomenon whereby if an individual is selected on the basis of having an extreme measurement value at time point one, their second measurement value will, on average, be closer to the mean of the population distribution (due to the influences of measurement error and normal within-subject variation). A corollary of this effect is that \( t_2 \) severity is often a significant covariate of change in symptom score between \( t_1 \) and \( t_2 \) – meaning that individuals with higher initial scores may appear to show the greatest improvement in symptom levels at follow-up, even when the true magnitude of change does not vary across individuals (see 10 for a worked example). The fact the \( t_2 \) score is used to calculate both quantities (i.e. they are mathematically coupled) results in further inflation of this relationship (see 20).

Care should therefore be taken when key predictors in response algorithms closely index \( t_2 \) severity, as this may result in a poorly generalising model. However, in previous studies in psychiatric datasets, baseline severity score is usually included among the features used to train response prediction algorithms (e.g. 21–23).

These factors may help explain why previous attempts to apply machine learning approaches to outcome prediction in psychiatric datasets have thus far had limited success in terms of out-of-sample (unseen data) classification. For example, a recent methodologically rigorous trial aiming to predict significant response (remission) following treatment with a particular antidepressant achieved only ~60% classification accuracy when the model was applied in external validation datasets. However, as previously noted, tools with only modest true predictive value may still have reasonably high clinical utility compared to current best practice; therefore this is still an approach very much worth pursuing.

Potential solutions

Clinical trial design
The problem of identifying true response heterogeneity is a problem of appropriately partitioning variance components in observed outcomes. The ability to properly identify differential response to a particular treatment in different individuals requires replication at the level at which the differential response is claimed (i.e., that particular treatment in that particular individual). Differential treatment response (i.e. identification of patient by treatment interactions) can therefore be identified by use of repeated period cross-over designs – a form of trial where each participant receives both placebo and active treatments more than once. However, in practice, these designs are rare, as they are likely to be impractical (prohibitively lengthy and expensive) and/or unethical. This kind of design also assumes that treatments wash out fully between administrations, which might not be reasonable for some interventions (e.g. psychological therapies).

Training data definition and selection
An alternative approach is to improve the way data from existing RCTs is used to train predictive models. For example, it has been suggested that the uncertainty in each individual’s ‘response’ (change in symptom score in the active treatment group) could be expressed as a confidence interval by reference to the standard deviation of the change scores in the control (placebo) group multiplied by the appropriate value from the \( t \) distribution (e.g. individual change score \( \pm 1.96 \times SD \) of control arm changes for a 95% CI, see 24). The probability that any given individual in the intervention group is a true responder (true change score is greater than the minimum clinically significant change) can then be derived from individual CIs using a Bayesian approach. Appropriate supervised learning algorithms could then be trained to predict (continuous) treatment response probability, as opposed to dividing individuals into binary response categories (e.g. using Gaussian process regression).
It also may be important to think carefully about the nature of the predictors (features) included in supervised learning models trained data – as those that reference initial clinical severity may be vulnerable to regression to the mean-related artefacts. There are statistical methods that have proposed to correct for regression to the mean when correlating t_{ij} symptom changes with initial severity level (see 20). However, these may require additional measurements (e.g. multiple estimates of t_{ij} value, in order to control for effects of measurement error).

**Counterfactual probabilistic modelling**

When a particular experiment is not feasible, one alternative is to train models from observational (non-experimental) data that are able to make counterfactual predictions – i.e. of the outcomes that would have been observed, had we run that particular experiment. For example, Saria and colleagues have recently developed a counterfactual Gaussian process (CGP) approach to modelling clinical outcome data26. The CGP is trained on observational (non-experimental) time-series data, in order to form a model of clinical outcomes under a series of treatments in continuous time. Crucially, the CGP is trained using a joint maximum likelihood objective, which parses dependencies between observed actions (e.g. treatments) and outcomes in the data. This feature allows prediction of how future trajectories (symptom levels) may change in response to different treatment interventions, and has previously been shown to successfully predict real clinical data (renal health markers following different kinds of dialysis,20,27).

This modelling approach requires datasets with semi-continuous measurement of the relevant clinical outcome (both pre- and post-treatment), in order to generate hypothetical treatment response traces – a kind of data that is not usually available from existing RCTs. Given sufficient attention to patient confidentiality and other ethical concerns, it may be possible to obtain appropriate training data from health service clinical records; however, frequency and consistency of symptom reporting may pose analytical problems (e.g. 27). The use of personal devices such as smartphones or other wearable technology to regularly self-record symptom levels may be a potential source of this kind of data in the future, given sufficient insight and patient compliance (e.g. 28). The CGP approach also rests on two key mathematical assumptions: that there will be a consistency of outcomes between training observations and future outcomes, given a particular treatment; and that there are no important confounding variables missing from the dataset26. It may require careful consideration as to whether these are reasonable assumptions for modelling psychological and chronic pain symptomatology.

**Conclusions**

The issues discussed above underline the importance of focusing on where data comes from when considering strategies for personalised medicine. In particular, it is problematic to designate individual data points from a conventional RCT design as ‘responders’ or ‘non-responders’ to a particular treatment, as this is in effect a single-arm (no control) study not adjusted for other important sources of response variation. This might be particularly important when considering patients with episodic, chronically-relapsing disorders as control variability is likely to be high (and symptom measurement itself is often imprecise). One solution to this problem is to use data derived from repeated cross-over design clinical trials, although in practice these can be prohibitively difficult and/or ethically problematic. It may be possible to alleviate these issues with careful model design, but this may still require changes to the way data is collected and monitored in the future in order to maximise potential clinical utility.

**Competing interests**

No competing interests were disclosed.

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**References**


This article represents a valuable contribution to the developing literature on quantification of individual responses to treatments and identification of patients who respond positively to treatments. Perhaps there should be some attention to the issue of identifying negative responders, since it is more important to avoid harming individual patients than to miss out on benefitting them. By "harm" I don't mean side effects. I also point out in my review that prediction models could be developed from identified modifiers of the treatment effect in controlled trials, something that you should also mention. I also have a strong view about repeatability of individual responses to treatments, something that you will need to address by refuting my claim or accommodating it. Otherwise there are only minor points for you to consider. These were all made on a first and only read-through, so although you explain some points further on, it is important to avoid any confusion in the first place.

"This may be particularly problematic in the case of psychiatric and chronic pain symptom data, where both within-subject variability and measurement error are likely to be high." I don't understand inclusion of "within-subject variability"? Are you referring to real changes of individual subjects pre to post the treatment that occur even in the absence of an active treatment? I normally think about that as another kind of measurement error. Perhaps you need to clarify by stating "both within-subject variability over the period of the treatment and short-term measurement error". Also, solitary "this" is a grammatical error known as an ambiguous antecedent. There are a few other instances in the manuscript that need to be fixed.

"especially in arenas where the potential clinical benefit is so large" I don't understand the use of "so". Are you referring to the arenas of psych disorders and chronic pain? Are the "potential clinical benefits" in these arenas any larger than in any other arenas of pathology? And you haven't established (in the Abstract, anyway) that there is the potential for large benefit when individual responders have been characterised. Maybe something along the lines of "trustworthy identification of characteristics of positive responders to treatments could result in substantial clinical benefit in psychological disorders, chronic pain, and other pathologies."

In the Introduction, you make it clear–at some length–that non-responders to one kind of treatment could be responders to another, and it may therefore be possible to improve the health of the majority of patients by targeting specific patients with specific treatments, where the pathology has a range of underlying causes and a range of treatments is available. Maybe you need to make that clearer in the Abstract.
“variables that may potentially relate” is a “double doubtful.” Remove either “may” or “potentially”.

“i.e. the reliability of distinguishing between treatment ‘responders’ and ‘non-responders’”. I think validity would be a better word than reliability. Also no need for quote marks.

e.g. and i.e. normally have commas after them and are used only in parentheses.

“Crucially, we can only draw this inference by direct comparison...” Make it “Crucially, we can draw this inference only by direct comparison...” Check for any other instances of this misplaced modifier.

You tend to use too many parenthetical asides. Remove parentheses from as many as you can.

"Formally, to properly infer whether a particular participant responded or didn’t respond to a particular treatment, we would require knowledge of what would have happened if a key event (treatment administration) both did and did not occur (a form of counterfactual reasoning), which is not possible in the real world." I found this sentence confusing. What is counterfactual reasoning? If I understand this sentence correctly, I disagree with it. In crossovers it is possible to determine the outcome with an individual who received the active and the control treatment. You go on to state that yourself.

"(e.g. time of day, stress level, etc.)" Either e.g. or etc., but not both!

"Measurement error may be higher than [in] other areas of medicine, as the main tools used to assess clinical outcomes are patient or clinician-completed questionnaire measures, which are relatively low[-]precision tools." I think this statement is false, so you’d better support it with references. The square root of the alpha reliability (which provides an upper limit to the criterion validity correlation of multi-item instruments) and short-term retest reliability ICC could well be high enough to reasonably identify responders to short-term treatments. Even VASs have high short-term ICCs. The ICCs over periods for long-term treatments are likely to be a different story, as you point out.

"If the variation in outcome due to these sources is greater than that due to any true individual differences in treatment response, it will be very hard to detect the latter under a conventional RCT framework.” It depends what you mean by "detect". To be on the safe side, perhaps you should state “The greater the variation in outcome due to these sources compared with that of true individual responses, the harder it will be to characterize the latter in a controlled trial.” Note that I have removed superfluous words. Bottom line is that you can make up for the short- and log-term errors with a big-enough sample size, at least for characterizing the mean effect and its modifiers.

“The fact the t1 score is used to calculate both quantities...” I fully understand regression to the mean, but I re-read this paragraph and still don’t know what "both quantities" refers to.

"The ability to properly identify differential response to a particular treatment in different individuals requires replication at the level at which the differential response is claimed (i.e., that particular treatment in that particular individual).” This rather obscurely worded claim has been made by others, but it is false. You can have a patient who responds individually to a treatment on one administration of the treatment. Whether that patient would respond similarly again following washout and reapplication of the treatment is irrelevant. What matters is that the patient has obtained benefit from the treatment when it was applied the first time. Period. Whether you can adequately quantify the extent of an individual patient’s response to the (first) application of the treatment depends on short- and long-term errors of measurement and on the magnitude of the response in that patient, but regardless, you can certainly
characterise modifiers of the treatment effect with realistic sample sizes: 4x the sample size required to characterize the mean effect (Hopkins, 2006). The identified modifiers could then be used to build courses-for-horses (treatments-for-patients) prediction models, something you haven't considered. Anyway, there is no need to confuse everyone by raising the spectre of repeatability of the treatment effect in individuals.

"However, these may require additional measurements (e.g. multiple estimates of t1 value, in order to control for effects of measurement error)." No, repeating the t1 assessment reduces but does not eliminate the effect of regression to the mean. What you need for the adjustment is the reliability ICC over the time-frame of the treatment. In fact, the control group effectively provides that: when you predict the likelihood of an individual's response in a controlled trial using a mixed model in which the pre-test (t1) is included as a modifying covariate, you have controlled for (adjusted away the effect of) regression to the mean.

I don't understand the paragraph headed Counterfactual probabilistic modelling. You will have to explain what is going on here without the jargon, for my benefit and for those who are even less statistically savvy. I see the word "trajectory" in there, which suggests to me that you are talking about clinical trials with multiple repeated measurements during the course of the treatment. That's a luxury that may not be available in many settings, and in any case, it requires an appropriate model for the time course, which is bound to be non-linear. You still need a control group, if you want to eliminate the contribution of the placebo effect.

Ah, I see you provide some explanation in the next paragraph. Please make the preceding paragraph clearer.

"The CGP approach also rests on two key mathematical assumptions: that there will be a consistency of outcomes between training observations and future outcomes, given a particular treatment…" No. See above.

"and that there are no important confounding variables missing from the dataset." Be more explicit. If it's a properly balanced controlled trial (or randomized with a sufficiently large sample size), what's the problem?

"One solution to this problem is to use data derived from repeated cross-over design clinical trials…" Well, no, because it introduces what I called above the spectre of repeatability.

"It may be possible to alleviate these issues with careful model design…" Explain "careful". You will need to have identified the point(s) explaining "careful" previously, as this sentence is in the Conclusions.


Is the topic of the opinion article discussed accurately in the context of the current literature? Yes

Are all factual statements correct and adequately supported by citations? Partly

Are arguments sufficiently supported by evidence from the published literature?
Partly

Are the conclusions drawn balanced and justified on the basis of the presented arguments?
Partly

**Competing Interests:** No competing interests were disclosed.

**Referee Expertise:** Research design and analysis, with special reference to physical activity, sport and lifestyle.

I have read this submission. I believe that I have an appropriate level of expertise to confirm that it is of an acceptable scientific standard, however I have significant reservations, as outlined above.

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In this manuscript, the authors discussed the concept of inter-individual differences in response to treatment interventions, particularly those focussed on psychological-related outcomes. The consideration of inter-individual responses is an important issue and the authors provide further insights previously not considered in detail within the domain of psychology. The topic is generally discussed accurately in the context of the current literature, statements are generally correct and supported by relevant citations. I have thoroughly read and considered the manuscript, which was interesting in content and constructed with a logical flow. I have only minor comments for the authors’ consideration.

1. The article focuses on the prediction of response heterogeneity, especially prediction of responders/non-responders. Have you considered the roadmap that has been suggested¹ to actually confirm whether the general amount of true heterogeneity is clinically important or not BEFORE we might explore for predictors of that heterogeneity?

2. Page 3, Right hand Column, Lines 27 onwards– whilst discussing RCT trial design and highlighting that responders/non-responders cannot be identified through the analysis of intervention sample data alone, perhaps it might be appropriate to address any research making similar claims even in the total absence of any control sample data.

3. Page 4, Left hand column, Line 8 and conclusion. The arguments that you make on this point, particularly in the conclusion are, at present, unsupported by scientific literature and require justification. You allude to the fact that a lack of ‘true’ counterfactual information makes an RCT in effect a single-arm (no control study). It is agreed that one cannot say with 100% certainty whether the intervention group as a whole or any specific individual in the intervention group is a positive responder, as what would have happened to that person if they had been in the control group is of course unknown. This is the fundamental counterfactual basis of the RCT. Nevertheless, as the control group variability over the same time period as the intervention effectively provides our best
guess of the counterfactual (what would have happened to individuals in the intervention group if they had been in the control arm), I feel that this applies to changes at both the group mean and the individual level, and that disregarding RCTs as ‘single-arm studies’ is unsupported. According to the previously-mentioned “roadmap” that has been presented, the analysis of the control group changes (specifically the comparison of change variance between treatment and control) can provide information as to what the general clinical importance is of “true” individual response heterogeneity. By “true”, one knows from this comparison whether the overall amount of heterogeneity in changes surpasses the overall amount of random within-subject heterogeneity of changes in the control group. If heterogeneity of change is similar between treatment and control, it could be argued that moving on to attempts to predict treatment response variability is a somewhat meaningless exercise.

4. Page 4, Left hand column, Lines 43 – 54. Whilst discussing regression to the mean and the mathematical coupling of pre- to post change scores, the use of covariates (especially baseline values of the study outcome) in the statistical model (ANCOVA) could be suggested as a potential solution to this – a notable absence in many studies’ data analyses.

5. Pages 4 – 5. You make a number of pertinent suggestions for potential solutions to the problem, and briefly allude to the methods recently suggested \(^1\).\(^2\). We have suggested how this might be approached in RCTs and tied to an appropriate anchor usually a minimal clinically important difference or smallest worthwhile change \(^1\).\(^2\). Addressing these issues may assist the reader in applying this methodology in their applied practice and/or research environments.

References

Is the topic of the opinion article discussed accurately in the context of the current literature?
Yes

Are all factual statements correct and adequately supported by citations?
Partly

Are arguments sufficiently supported by evidence from the published literature?
Partly

Are the conclusions drawn balanced and justified on the basis of the presented arguments?
Partly

**Competing Interests:** No competing interests were disclosed.

We have read this submission. We believe that we have an appropriate level of expertise to confirm that it is of an acceptable scientific standard.
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