

Some reasons why preclinical studies of psychiatric disorders fail to translate: what can be rescued from the misunderstanding and misuse of animal 'models'?

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Abstract

The repeated failure of animal models to yield findings that translate into humans is a serious threat to the credibility of preclinical biomedical research. The use of animals in research that lacks translational validity is unacceptable in any ethical environment and so this problem needs urgent attention. To reproduce any human illness in animals is a serious challenge, but this is especially the case for psychiatric disorders. Yet, many authors do not hesitate to describe their findings as a 'model' of such a disorder. More cautious scientists describe the behavioural phenotype as 'disorder-like', without specifying the way(s) in which the abnormal behaviour could be regarded as being analogous to any of the diagnostic features of the disorder in question. By way of discussing these problems, this article focusses on common, but flawed, assumptions that pervade preclinical research of depression and antidepressants. Particular attention is given to the difference between putative 'models' of this illness and predictive screens for candidate drug treatments, which is evidently widely misunderstood. However, the problems highlighted in this article are generic and afflict research of all psychiatric disorders. This dire situation will be resolved only when funders and journal editors take action to ensure that researchers interpret their findings in a less ambitious, but more realistic, evidence-based way that would parallel changes in research of the cause(s), diagnosis and treatment of psychiatric problems in humans.

Background

Immense effort is being invested in devising remedies for the poor reproducibility of preclinical biomedical research. It goes without saying that the design of experiments, and the way they are carried out, are crucial for ensuring that the conclusions are valid. But, in terms of helping to ensure successful translation of preclinical research into humans, the campaign to improve reproducibility is merely tinkering around the edges unless the scientific rationale for an experiment is sound. Taking steps to improve the reproducibility of experimental results will not improve translation if the interpretation of the research findings is misleading. A failure to translate is not only a major setback for the field but, when animals have been used at any stage of the process, it is ethically unacceptable if the failure was avoidable.

There have been several recent commentaries on factors that contribute to failed translation. These have covered many important points, but have tended to focus on topics such as: species differences; methodology; experimental environment; neglected complications arising from the use of genetically-altered animals; subjective and systematic bias; skills deficits; and even a need to redefine the whole strategy for preclinical research (e.g.,¹⁻³). However, a fundamental problem, which has received surprisingly little attention, is that some procedures are being used as ‘models’ of complex human disorders when that was not their intended purpose and the underlying assumptions have never been validated⁴.

This article does not offer a detailed critique of the wide range of procedures that are used in research of the causes and treatments for psychiatric disorders, not least because that would be a task with almost unlimited scope. It is also not intended as a discourse of the important ethical and philosophical debates arising from species differences in neuroanatomy and neuronal networks, which lead some authors to conclude that the development of an animal model for any human brain disorder can never be a realistic objective^{5,6}. Instead, the baseline for this article is that not only do many scientists believe that such models exist already, but they are using them with impunity.

That would be acceptable if the assumptions that underpin the model(s) have been validated, and if it is acknowledged that their validity depends on the experimental context and the research objectives. The aim of this article is to highlight the extent to which those vital steps are typically ignored and, as a consequence, both the rationale for the experiments and the conclusions arising from the work risk being fundamentally flawed. These problems do not rest on the fine details of each ‘model’ or psychiatric disorder; they are generic and so this article draws on some striking

examples from preclinical research of depression and antidepressant in order to illustrate some ways in which validity can be compromised. Particular attention is devoted to an appraisal of the Forced Swim Test (FST) because that procedure has attracted a good deal of criticism and is the focus of a campaign to ban its use altogether.

All these 'models' have been used for many years and so it is realistic to assume that many scientists will continue to use them, as before, despite their limitations. However, if the problems discussed below are not resolved, there is a risk that confidence in the scientific merits of all procedures used in preclinical psychopharmacology will wither irrevocably, even though many have made invaluable contributions to the field.

To help prevent that from happening, the final section suggests remedial actions that could help to strengthen the validity of inferences from a program of research that incorporates the use of a 'model' of a psychiatric disorder or 'disorder-like' behaviour.

Reasons for studying behavioural phenotypes

There are two main reasons for studying the behaviour of laboratory animals in preclinical research of psychiatric disorders. One is to induce behavioural and / or biological responses in the animals that are analogous to the human disorder of interest: i.e., to produce a 'model' of at least one aspect of the illness. The neuronal mechanisms underlying these responses can then be interrogated in more detail, which could point the way to new and better treatments. A problem arising from this type of research is that the behavioural phenotype produced by the procedure is often assumed to be analogous to the human disorder, even though it does not begin to qualify for such a 'diagnosis'.

The other reason for studying animals' behaviour is to test whether or not a potential new treatment merits further development for a given clinical indication. This involves evaluation of the effect of the candidate treatment on the behaviour of an animal during exposure to a standard test procedure: e.g., the Forced Swim Test or the Tail Suspension Test. The important point about these procedures is that none requires the baseline behaviour (i.e., the behaviour of untreated animals) to be a model of any aspect of the target illness. All that is needed is for all treatments of the same class, with confirmed therapeutic efficacy in humans, to produce a clear and consistent change in any aspect of the animals' behaviour. That response then acquires '*predictive validity*' and can be used to screen new candidates for their potential as effective treatments in humans. However, a common misunderstanding, which is discussed in more detail,

below, is that it is often assumed that the behaviour that is being evaluated is a model of psychiatric disorder, or disorder-like behaviour, and that its prevention by a test treatment indicates that it has ‘cured’ the illness. This is a particularly common problem when these procedures are used to look for changes in the behaviour of animals that have been genetically altered in a way that is thought to be relevant to the psychiatric disorder of interest.

The challenges for animal ‘models’ of a human psychiatric disorder

As the publication archive confirms, many researchers believe that it is plausible to claim that an experimental intervention has induced an animal ‘model’ of a human illness when there is clear pathology or biomarker, which is common to both species: e.g., when animals have a specific type of cancer or a microbial infection. But even in those cases, translation into humans and development of effective treatments has not been as straightforward as expected. It is far more difficult to be confident about the validity of an animal model when the human disorder comprises several different contributory factors (e.g., hypertension or metabolic syndrome). The development of animal models of psychiatric disorders is especially challenging because no defining pathology or biomarkers for any of these have been discovered so far.

There are several different systems to steer the diagnoses of psychiatric illnesses (e.g.,^{7,8}). Although the details of the criteria that qualify for a formal diagnosis differ slightly from one to another, the qualitative aspects of symptoms and signs that are regarded as common in depressed humans are listed in Table 1.

Some of these features are self-evidently analogous in humans and other animals and their evaluation is fairly straightforward; this would be the case for monitoring changes in body weight, disruption of sleep architecture or feeding patterns, for instance. Similarly, cognitive impairments that are arguably equivalent to those expressed by humans can also be evaluated in animals, using tests to monitor their spatial memory, discrimination of novel aspects of their environment, focused attention and so on. By contrast, the equivalence of abnormalities expressed in animals and other aspects of depression is less clear-cut.

For instance, the loss of rodents’ innate preference for sweet fluids, after experience of a prolonged series of mild stressors, is widely regarded as a sign of ‘anhedonia’, which is a prominent feature of depression. However, it should be borne in mind that the detailed DSM-5 criterion for anhedonia in depression is ‘*a markedly diminished interest or pleasure in many, or all, activities, nearly every day*’: i.e., the anhedonia is not confined to a gustatory preference.

Similarly, rodents' submissive behaviour after social defeat is often described as 'depression-like' behaviour, but it is not clear what aspect of depression is being emulated. Low self-confidence is a remote possibility even though, in depressed humans, that symptom is not normally attributed to threats of physical attack from a dominant conspecific.

In neither of those cases, is there any evidence to indicate the animals' state of mind and so the description of the behavioural abnormality as being 'depression-like' is based on risky anthropomorphic assumptions. That limitation taps into the major obstacle for modelling any psychiatric disorder, which is that many of the key diagnostic criteria are subjective and cannot be evaluated in animals at all (e.g., suicidal ideation in depression; flashbacks and survivor guilt in Post-Traumatic Stress Disorder; hallucinations in schizophrenia).

To add to these problems, a diagnosis of a psychiatric disorder rests on patients expressing combinations of symptoms and signs, which are long-lasting (often, life-long). Some are given more weight than others and their number and type can differ substantially from one patient to another and yet still qualify for a diagnosis of depression. Moreover, the severity of the illness can wax and wane, with periods of remission that have unpredictable duration. Another confounder is that some symptoms and signs are evident in more than one psychiatric disorder. Given that many disorders are often comorbid and that some of the features of comorbid illnesses overlap, it can be difficult, if not impossible, to attribute any particular behavioural abnormality to either one, alone.

It is reasonable to expect the criteria for a valid model of depression in animals to be equivalent to those required for a diagnosis of depression in humans. However, DSM-5, for instance, requires patients to express at least five of nine symptoms for at least two weeks. Even if the bar for a diagnosis in animals is lowered as much as possible, by ignoring important details of each feature that psychiatrists would look for in humans (e.g. its duration – as in Table 1), it is clear that no experimental intervention (e.g., drug treatment, genetic alteration, neuronal lesion) has yet produced a collection of abnormalities in animals that would qualify as a 'model' of depression in terms of its symptom profile, let alone its severity and temporal instability.

To wriggle out of that problem, many authors resort to describing the behavioural abnormality they have studied as 'depression-like' instead, although it is never made clear what 'depression-like' actually means (c.f. Table 1; see also:¹). Nevertheless, this is a promising approach because current research strategies are moving towards the study of endophenotypes, whereby a specific abnormality can be linked with an underlying biological process: such as a neuronal network or

genetic mutation⁹. Even so, if it is to be claimed that an endophenotype is analogous in humans and other animals, its description still needs to be far more specific than merely 'depression-like'.

Some procedures that are used to produce animal 'models' of depression

This section describes some important examples of procedures that are used as preclinical of depression in rodents and outlines their strengths, limitations and common pitfalls.

Reserpine

This was the first animal 'model' of depression. Reserpine is a naturally-occurring alkaloid that was used to treat hypertension and mania until around the early 1960s. At that time, it was asserted that this drug caused suicidal depression a few patients (c. 12-15%), but that now seems unlikely¹⁰. Nevertheless, that belief provided a rationale for studying the effects of reserpine in animals on the grounds that it would induce depression in other species too. This assumption was further encouraged by evidence that a new class of antidepressant drugs, which had just been discovered (the 'tricyclics'), prevented reserpine-induced hypothermia¹¹. The finding that the tricyclics had negligible effect on the behavioural response to reserpine treatment (akinesia) was basically ignored.

As well as inducing hypothermia and hypotension, reserpine also depletes monoamine-releasing neurons of their pool of neurotransmitter (norepinephrine, serotonin or dopamine). This action seemed to strengthen the endorsement of the model because, at that time, it was thought that depression was caused by a deficit in monoamine transmission. That is no longer the case and, given that none of these abnormalities (hypothermia, hypotension or depletion of brain monoamines) are evident in depressed humans, reserpine treatment is now generally deprecated as a model of depression. Nevertheless, it is still used by some laboratories to study "reserpine-induced depression" [*sic*] and to screen novel compounds for their potential as antidepressants, even though prevention of the hypothermia only ever seemed to happen with drugs that block neuronal reuptake of noradrenaline¹². Examples can be found on PubMed using the keywords: reserpine / depression / model.

Learned Helplessness

This procedure involves exposing animals to uncontrollable, unpredictable stress (bouts of electric footshocks) in an environment from which they cannot escape. The animals freeze on subsequent exposure to that environment, even when given the opportunity to escape (see:¹³): i.e., they develop an 'escape deficit'. When first discovered, this behaviour was somewhat arbitrarily called 'learned helplessness'. That label strongly implied that the stress induces a particular state of mind in the animals, which is analogous to the feeling of helplessness, experienced by depressed humans. That implication was supported to the extent that the animals also show impaired grooming, loss of appetite and cognitive deficits, which are arguably analogous to features of depression in humans (c.f., Table 1). The finding that all these changes are prevented by treatment with antidepressants¹⁴ further strengthened that possibility.

Nevertheless, the question of whether or not learned helplessness is analogous to depression in humans has been debated continually over the last 50 years. It has even been suggested that it is a model of post-traumatic stress disorder (PTSD), rather than depression¹⁵. The prevailing view is that the escape deficit arises from an innate, default passivity, rather than 'depression', which is triggered by a lack of control over environmental stressors. A further suggestion is that finding ways of preventing this response (behavioural passivity) could hold the key to antidepressant¹⁶.

The debate over learned helplessness illustrates how it can be really difficult to be confident about what human state an abnormal behaviour is actually being emulated by the 'model'. That setback, together with concerns about the severity of this procedure, means that it is ethically controversial, which probably explains why it is rarely used in jurisdictions with a strong emphasis on the 3Rs and welfare of laboratory animals.

Olfactory bulbectomy

This is the best validated model of depression and involves bilateral surgical removal of the olfactory bulbs in rodents. Notwithstanding species differences in the effects of bulbectomy on animals' behaviour and physiology¹⁷, this procedure causes a wide range of abnormalities in the animals, many of which echo what happens in humans suffering from depression. These involve not only changes in neuronal networks that are consistent with our understanding of the neurobiology of depression^{18,19}, but also disruption of the immune (cytokine production) and endocrine (raised plasma cortisol) systems, in ways that are common in depressed patients. A strong piece of evidence that distinguishes olfactory bulbectomy from other models is that many

of these changes are prevented by prolonged, but not short-term, treatment with an antidepressant drug²⁰ (c.f., below).

This diverse evidence offers a good example of how validation of a putative animal model involves confirmation that a wide range of abnormalities seen in depressed humans are reproduced in animals too; these include neuroendocrine, immune, therapeutic profile and timescale, not merely a change in a single aspect of their behaviour. By contrast, most rodent 'models of depression rely merely on comparisons of the change in their motor behaviour after an experimental challenge (e.g., when suspended upside down or immersed in a bucket of water) and do not consider the other physiological changes that are often associated with the human disorder.

Despite the strengths of this model, it does have some limitations, which are widely acknowledged. One is that olfactory bulbectomized animals are hyperactive, which is the opposite of what typically happens in depressed patients. Another is that depression in humans clearly does not involve physical ablation or section of the olfactory bulbs. However, there are reports that the sense of smell is impaired in some depressed patients²¹, which raises the fascinating possibility that a functional deficit in this brain region could explain this subgroup of patients. Finally, the surgical intervention needed for olfactory bulbectomy is regarded as a severe procedure under the European legislation and so its use needs stringent ethical justification.

Chronic mild (unpredictable) stress (CMS / CUMS)

As mentioned above, rodents normally have a strong preference for sweet drinks, but this is diminished after experience of a series of mild stressors. The stressors do not need to be noxious or cause overt physical discomfort, but merely involve disrupting animals' environment or daily routine. When used as intended when it was first developed in the UK, the procedures apply stressors such as: short-term food or water restriction; continuous lighting, cage tilt, group-housing, wet bedding, low temperature (10°C), intermittent white noise (85dB); novel object in cage; strange odour (air freshener). The animals experience one of these stressors each day, for several weeks, after which they develop a deficit in the Sucrose Preference Test²².

Because the loss of sucrose preference after CMS is prevented by antidepressants, it is widely assumed that this change, alone, justifies the use of CMS to produce a model of depression, or depression-like behaviour. A more realistic and specific description would be that this deficit is equivalent to a loss of the rodents' motivation to experience pleasure, which is arguably relevant

to 'anhedonia' in depressed humans (but see above). Whether or not this is the case, it should be borne in mind that anhedonia is a common element of other psychiatric disorders too (e.g., schizophrenia, bipolar disorder, obesity and PTSD).

A particular concern about the CMS / CUMS procedure is the variability in the types of stressors used in different laboratories. In some cases, the protocol involves social isolation in combination with stressors such as: 24 h food deprivation and/or 24 h water deprivation; forced swimming in ice-cold water (4-6°C); placement in an oven (45°C), several hours of physical restraint; several hours of continuous loud white noise. The animals can experience at least one or two of these stressors, every day, for several weeks. Recent examples of these experiments can be found in PubMed using keywords: stress / chronic / mild / sucrose.

It is hard to understand how any of these types of stressors could be regarded as 'mild', still less so when the cumulative harm to the animals is taken into account. As well as the ethical considerations arising from these studies, it is important to bear in mind that neuronal responses to stress are strongly dependent on the type, duration, frequency and severity of the stress (reviewed by: ²³). It cannot be assumed that findings that emerge from one protocol are typical, or that studies using different protocols can be compared in any meaningful way. It is also worth considering how these types of stress could be relevant to those that can trigger depression in humans, which often involve loss of control (e.g. bereavement or redundancy), as opposed to PTSD or other long-term consequences of experiencing physically traumatic, life-threatening stress.

Social defeat stress

There are several different types of tests in this category, but they all evaluate changes in animals' behaviour after a social challenge between dominant and subordinate animals. Following several bouts of social defeat, the subordinate (defeated) animal typically develops a loss of sucrose preference, a deficit in grooming, reduced body weight and disruption of sleep architecture (Table 1)^{24,25}. On that basis, this is widely described as a model of 'depression-like' behaviour. However, that inference is somewhat undermined by the gamut of research that has used this procedure to study 'anxiety' (e.g.,^{26,27}).

As with CUMS procedures, a major concern about these studies is that there are appreciable differences in the protocols used in different laboratories, particularly in respect of the time for which the animals are left to interact. In some laboratories, as in the UK, the animals are separated

immediately after the first contact, but retain sight and smell of each other²⁸. Elsewhere, the physical interaction between the animals is scored over periods that can last up to 30 min each day. The different versions of these tests have implications not only for the interpretation of the findings, but also for 3Rs compliance, animal welfare and what qualifies as an ethically acceptable procedure in different jurisdictions.

Predictive drug screens for antidepressants are not models of depression

A second cluster of procedures comprises those that are used as fast throughput, predictive screens to test novel compounds at an early stage of their development. To achieve the status of a predictive screen, all drugs of a given class must affect an aspect of the animals' behaviour in a specific way. For instance, most drugs that induce the characteristic behavioural 'serotonin syndrome' in rodents, which includes head-twitches, 'wet-dog shakes' and Straub tail, induce hallucinations in humans. Although none of these behaviours is seen in humans (especially Straub tail!), there is a well-justified expectation that if a test drug induces this cluster of abnormal behaviours in rodents, there is a high risk that it will induce hallucinations in humans (see²⁹).

Predictive screens for antidepressants have less pharmacological precision because these drugs interact with many different types of molecular targets in the brain. These range from neurotransmitter receptors and transporters to genes that influence glial cell connectivity. Nevertheless, as discussed below, the behavioural response to certain experimental procedures is modified in a consistent way by all antidepressants. In such cases, the effect of a test drug on the animals' behaviour response can be an invaluable guide to whether or not it is likely to act as an antidepressant in humans.

Some of the procedures, described above are used in this way (e.g. CUMS and social defeat). Others include the Tail-Suspension Test (for mice) and various measures of impaired episodic memory (e.g. novel object or novel location recognition). However, one test that is widely used for this purpose is the Forced Swim Test; this is discussed in detail, below, because it is an excellent example of how the validity of an experimental procedure can depend on the experimental objectives: i.e., the FST can be valid for one purpose (as a predictive drug screen) but not another (a model of depression).

The Forced Swim Test (FST)

This procedure involves immersing rodents in a tank of water from which they cannot escape. After a few minutes, the animals stop swimming and adopt a posture ('immobility'), which enables them to float with their noses held above the surface of the water. All established antidepressant drugs increase the latency to adopt this immobile posture (e.g.,³⁰) and this test has been used as a predictive screen for antidepressants for over 40 years. It is striking that even S(+)-ketamine, which is a stereoisomer of the anaesthetic, ketamine, reduces immobility in the FST³¹ and was given FDA approval, in 2019, as a lead compound for a completely new class of fast-acting antidepressants.

One criticism of the FST points to evidence that many factors can affect immobility when the animals are tested in a drug-free state. It is inferred that the FST must be intrinsically unreliable because the variables that affect animals' baseline behaviour in this test will vary in different laboratories (e.g.,^{32,33}). However, the fact that all established antidepressants reduced immobility in this test, despite this baseline variability, actually strengthens, rather than undermines, its value as a fast throughput predictive screen for therapeutic efficacy in humans: i.e., the antidepressant 'signal still shows up above the baseline 'noise'.

Obviously, we cannot know how many compounds that would have made effective antidepressants returned a false negative result in the FST and so were not developed for the clinic. On the flip-side, the possibility that there are true false positives (i.e., drugs that reduce immobility in this test but do not appear as antidepressants in the clinic) cannot be ruled out either, but the strident criticisms of the FST have focused on specific examples. These need to be addressed because there are many reasons why a compound that looked promising in the FST might not reach the clinic and a verdict of a false positive finding in the FST might not be as clear-cut as is often claimed.

First, the estimation of the optimal dose in humans used to test for efficacy in Phase III clinical trials, informed by anisotropic estimates from animal studies, might have been incorrect: a higher dose might have turned out to be efficacious (e.g.,³⁴). Secondly, the drug might not have met safety requirements, or it had undesirable side-effects: i.e., its development stumbled after Phase I or Phase II and efficacy was never tested in the clinic. Thirdly, the drug simply turned out to be highly effective at treating a different disorder with a more promising market niche: e.g., sibutramine, marketed as an anti-obesity agent instead, and amoxapine, which was renamed atomoxetine and is now used to treat Attention Deficit Hyperactivity Disorder.

A specific group of compounds that has been highlighted as producing a false positive result in the FST include antihistamines. This is a particularly interesting example because many

antidepressant drugs are ligands for histamine receptors. In fact, an entire class of antidepressants, known as ‘tricyclics’, are molecular derivatives of the prototypical antihistamine, promazine and are antagonists of H₁ receptors. It is possible that antihistamines would have useful antidepressant effects in humans, were it not for the problem that H₁ receptor antagonists are highly sedative if they reach the brain, as anyone who suffers from hay fever will testify. Indeed, drugs that selectively target other types of histamine receptors are currently being investigated for their potential as antidepressants

Amphetamine and caffeine are also claimed to be convincing examples of drugs that produce false positives in the FST. Apart from the problem that amphetamine is highly addictive and so has restricted licensed approval (for treatment of Attention Deficit Hyperactivity Disorder), that criticism disregards the fact that it was the first drug treatment for depression and was used for over 30 years, until the 1960s, when the first antidepressants were discovered. It was regarded as being especially beneficial for patients expressing profound fatigue³⁵. Recent studies, using meta-analysis, continue to support the view that amphetamine can have beneficial effects in depression³⁶. Similarly, there is mounting evidence that caffeine³⁷ could make a useful adjunct to established antidepressant drug regimens³⁸. In short, neither of these compounds are false positives.

A final reason why some compounds that reduced immobility in the FST have not turned up as antidepressants in the clinic is that the reduction in animals’ immobility in the FST could actually be attributed to a non-specific increase in animals’ motor activity, rather than a reduction in swim-stress induced immobility (which is completely different), but that was not checked. Examples of this possibility include: green tea³⁹, ghrelin⁴⁰, probiotic supplements⁴¹ and NK₁ receptor antagonists. Evidence that a lack of functional NK₁ receptors causes locomotor hyperactivity in mice did not come to light until several years after the development of NK₁ receptor antagonists as antidepressants had been abandoned⁴².

However, the main problem with the FST is the widespread belief that, if antidepressants delay the onset of immobility of rodents, then immobility must be a ‘model’ of depression in humans. There is no evidence to support that assumption, which is an example of what Garner has called ‘a logical trap’¹. This misunderstanding seems to derive from early publications describing the FST in which the authors stated that: “*We suggested that this characteristic and readily identifiable behavioural immobility reflects a state of despair in the rat*” and they went on to describe the immobility as “*a behavioral state resembling depression*”⁴³. A later paper by the same group speculated that “*the immobile behaviour may reflect a state of lowered mood in the*

animal"⁴⁴. Even by 2001, this position had not changed: *"It was hypothesized that immobility reflected the animals' having learned that escape was impossible and their having given up hope. Immobility was therefore given the name 'behavioral despair' "*⁴⁵. It is clear that there was no evidence that immobility in the FST has anything to do with depression and that remains the case.

The possibility that the immobility is analogous to some aspect of depression is most unlikely, not least because depression is a chronic, relapsing disorder, whereas any change in the animals' mood while immersed in water would presumably dissipate soon after they are returned to their home cage. Also, the immobility is prevented by short-term treatment with an antidepressant drug (within 24h of the test), whereas the therapeutic lag for antidepressants in humans is typically 6-8 weeks, or more. These points, alone, are sufficient to rule out the immobility in the FST as a model of depression or even 'depression-like' behaviour.

Others share this skepticism and have suggested that the immobility is actually a passive coping (survival) mechanism³⁸, or that the immobility is driven by anxiety, not depression⁴⁶. Whatever the case, in discussing the use of the FST as a model of depression, and the failure of certain findings to translate into humans, Anyan and Amir⁴⁷ hit the nail on the head: *"There are two possible explanations for the discrepancies between human and animal research: either the underlying mechanisms driving depression and anxiety are distinct in humans and rodents or we are misinterpreting animal behavior. We believe the underlying mechanisms are conserved and therefore it is more likely due to interpretation error."*

It is interesting to speculate that immobility in the FST could be analogous to the psychomotor retardation seen in depression. If so, perhaps that is exacerbated by (swim) stress? There is evidence from one early study, which tested the psychomotor speed of depressed patients and found this was reduced when the task was made more stressful⁴⁸. Until this is confirmed, the only safe conclusion is that the FST enables measurement of 'stress-induced immobility'. To describe it as anything else, or to align it with any aspect of depression, is mere conjecture. In short, the merits of the FST as a useful, high-throughput predictive screen for antidepressant drugs is borne out by more than 40 years of evidence, but there is no reason to suppose that it produces an animal 'model' of depression. Indeed, it is not obvious what aspect of depression would be induced, or exacerbated, by immersion in a tank of water for a few minutes.

The Open Field as a follow-up (secondary) drug screen

As discussed above, if a drug is claimed to have beneficial effects on mood on the basis that it increases animals motor activity in procedures such as the FST, then it is essential to rule out the

possibility that these drugs are merely increasing animals' motor activity, nonspecifically. To achieve that, most studies go on to check the effect of the test drug on animals' ambulation in an arena, known as an Open Field. The Open Field Test is alluringly simple to carry out because it merely involves placing the animals in an arena for a few minutes and monitoring how much it moves around: apparently, neither the animals nor the experimenters need any expert training. However, there are many reasons why evaluation of locomotor activity in an Open Field is far from straightforward (see: ⁵¹).

The first, and most important, point to bear in mind is that the Open Field Test was developed to study animals' emotionality - not motor activity. However, it soon became clear that evaluation of differences in measures of animals' emotionality is confounded by differences in their ambulation in the arena and vice versa⁵². Secondly, animals' behaviour in this test depends on the physical characteristics of the arena, including its size, shape, the material used for its construction, light intensity and so on (see: ⁵³). Thirdly, animals express many types of behaviour in the Open Field and some of these will be incompatible with ambulation. For instance, a change in grooming or rearing, following a drug treatment or genetic mutation, will inevitably affect the animals' locomotor activity, indirectly, because they cannot express those behaviours and move around at the same time^{51,53-55}.

It follows that locomotor activity in the Open Field is not a single, independent experimental variable. As with all tests of this sort, changes in animals' behaviour in the Open Field are uninterpretable unless all other aspects of the behavioural profile have been taken into account. This is especially the case when the drug or genetic mutation being tested is intended to affect the mood of the animals. The best way to evaluate the effects of a drug on locomotor activity, specifically, is to monitor its effects on behaviour while the animals are in a familiar, naturalistic environment – ideally their home cage, ideally the animals' home cage⁵⁶ - and there is now equipment that makes that feasible.

What can be rescued from the misunderstanding and misuse of animal 'models'?

One of the recurrent themes of this article is that, despite there being no validated rodent model for full-blown depression, some protocols might induce behavioural change(s) that could be analogous to specific element(s) of that disorder. In such cases, authors should avoid using flimsy descriptions of what they have studied ('disorder-like') and should specify and scientifically justify the aspect(s) of the disorder they have in mind (e.g., anhedonia, low self-esteem, psychomotor

retardation). Of course, such specific inferences also need careful validation, together with an acknowledgment that they are not necessarily confined to a single psychiatric disorder.

That change of mindset would align closely with the growing interest in how human endophenotypes map onto specific neuronal pathways and/or genetic polymorphisms. This is the rationale for the international Prism Project (<https://prism-project.eu>). This program of research aims to gather data on social withdrawal and cognitive deficits, which are shared across multiple psychiatric disorders (Alzheimer's disease, schizophrenia and major depressive disorder) and to identify how the underlying biological parameters differ in these illnesses. One objective of this project is to discern distinct endophenotypes and then, through backtranslation, to develop valid, evidence-based animal models that would enable meaningful basic science research that is not permitted in humans⁹. This approach will be complemented, and informed, by parallel exploratory, investigation of new drug (IND) (pre-Phase 1, clinical trial) studies in humans, which is reducing the number of animals needed for approval of a full clinical trial, to some extent⁵⁷. A better understanding of the underlying endophenotypes is essential for personalized (stratified) medicine and to explain why some patients respond to a given drug treatment, while others do not.

This approach, which regards psychiatric disorders not as single, defined entities but as assemblies of abnormal endophenotypes⁹ is well advanced, but its basic principles have yet to permeate preclinical laboratories.

What can funders and journals do to help?

Both funders and journals are well placed to take steps that strengthen the rigor and validity of animal models: how they are used and how the findings are interpreted and described. Obviously, funders need to ensure the validity of the research they sponsor, but journals can help too. Yet, Editorial Boards have been remarkably nervous about taking any action to improve matters. An exception is the *Journal of Psychopharmacology*, which took the lead in 2019 by revising their Guidelines for Authors to include the point: “*Studies aimed at producing an animal ‘model’ of a psychiatric disorder (or ‘disorder-like’ behaviour) in humans should also include a statement to justify the extent to which the experimental procedure produces a validated animal analogue of the human condition, bearing in mind the diagnostic criteria specified in the Diagnostic and Statistical Manual of Mental Disorders (DSM-5®)*” (<https://journals.sagepub.com/author-instructions/JOP#Preclinical%20Studies>).

It is too early to assess how conscientiously authors are responding to this rubric, but it is certainly a nudge in the right direction. The adoption of a similar policy by other journals, which publish papers in which the authors claim to have used an animal model of a complex human disorder, is long overdue.

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Table 1 Core (qualitative) aspects of the symptoms and signs that contribute to a diagnosis of depression in humans and should also be expressed in animal ‘models’ of depression

| Diagnostic feature | | DSM-5 | ICD-10 |
|----------------------------------|--|--------------|---------------|
| <i>Psychological</i> | Sadness / depressed mood | ✓ | ✓ |
| | Diminished interest or pleasure in activities | ✓ | ✓ |
| | Cognitive impairment / poor concentration / indecisiveness | ✓ | ✓ |
| | Suicidal thoughts | ✓ | ✓ |
| | Feelings of worthlessness or (inappropriate) guilt | ✓ | |
| | Guilt or self-blame | | ✓ |
| | Low self-confidence | | ✓ |
| <i>Somatic vegetative</i> | Weight loss or gain | ✓ | |
| | Disturbed sleep (insomnia / hypersomnia) | ✓ | ✓ |
| | Fatigue / loss of energy | ✓ | ✓ |
| | Reduced or increase appetite | ✓ | ✓ |
| <i>Behavioural</i> | Agitation with reduction in physical movement | ✓ | ✓ |

