The place of implementation science in the translational medicine continuum

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There is a growing consensus that the transfer of knowledge from biomedical discoveries into patient and public benefit should be accelerated. At the same time there is a persistent lack of conceptual clarity about the precise nature of the phases of the translational continuum necessary to implement this. In this paper, we: (i) propose an integrated schema to understand the five sequential phases that link basic biomedical research with clinical science and implementation; (ii) discuss the nature of three blocks along this translational pathway; (iii) outline key issues that need to be addressed in removing such barriers. The five research phases described are: (0) basic science discovery; (1) early human studies; (2) early clinical trials; (3) late clinical trials; (4) implementation (which includes adoption in principle, early implementation and persistence of implementation). This schema also sets out three points at which communication blocks can occur. The application of ‘implementation science’ is in its early stages within mental health and psychiatric research. This paper therefore aims to develop a consistent terminology to understand the discovery, development, dissemination and implementation of new interventions. By better understanding the factors that promote or delay knowledge to flow across these blocks, we can accelerate progression along translational medicine pathways and so realize earlier patient benefit.

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Introduction

There is an emerging consensus that the transfer of knowledge from proven biomedical discoveries into patient and public benefit should be accelerated. At the same time there is little conceptual clarity, either about the precise nature of the phases of this ‘translational continuum’ or about the proper place within it of ‘implementation science’ (Eccles et al. 2009). In this paper, we aim to: (i) develop an integrated schema to understand the whole translational medicine continuum, consisting of five sequential phases that link basic biomedical research with clinical science and practice; (ii) discuss the nature of three important blocks between these phases; (iii) consider the place of implementation science within this continuum.

A schema for the translational medicine continuum

The best developed framework describing the development of new therapeutic interventions is that which refers to pharmacological drug discoveries. Such studies are divided into the five phases shown in the first row in Table 1. Subsequently, other schemes, which refer to non-pharmacological discoveries, have been elaborated (see Table 1). Within the UK, for example, a Framework for the Evaluation of Complex (largely psycho-social) Interventions has been described by the Medical Research Council using a similar sequence (Campbell et al. 2007; Craig et al. 2008). The National Institute for Health Research in England has established Biomedical Research Centres to support the conduct of translational medicine, which it sees as those investigations that begin with first-in-man studies and which continue up to, and including, early clinical trials (National Institute for Health Research, 2006). In parallel, within the USA a concerted scientific action programme has led to the National Institutes of Health Roadmap (Zerhouni, 2003), in which two ‘translational roadblocks’ have been described that delay knowledge transfer along the whole of the translational pathway (Zerhouni, 2005). Within the field of cancer research in the USA, for example, the President’s Cancer Panel has distinguished ‘early’ from ‘late’ translational studies (The President’s Cancer Panel, 2005). By combining
### Table 1. Five phases of the translational medicine continuum and three translational blocks

<table>
<thead>
<tr>
<th>Source</th>
<th>Phase 0 Basic science discovery</th>
<th>T1 Block</th>
<th>Phase 1 Early human trials</th>
<th>Phase 2 Early clinical trials</th>
<th>T2 Block</th>
<th>Phase 3 Late clinical trials</th>
<th>T3 Block</th>
<th>Phase 4 Implementation</th>
</tr>
</thead>
<tbody>
<tr>
<td>Drug discovery phases</td>
<td>Drug discovery: animal and laboratory/preclinical studies</td>
<td>Normal human volunteers: pharmacokinetics, safety and tolerability, determine safe dosage range and early side effects</td>
<td>Exploratory clinical studies: of efficacy in target population: patient volunteers, identify optimal doses, compare safety profiles with existing treatments, establish necessary treatment duration</td>
<td>Confirmatory clinical studies: of effectiveness and safety in target clinical population, identify less frequent and longer term side effects</td>
<td>Market launch and post-marketing surveillance</td>
<td></td>
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<td>UK: MRC framework for evaluation of complex interventions 2007 (Campbell et al. 2007) 2008 (Craig et al. 2008)</td>
<td>Pre-clinical [theory] explore relevant theory to ensure best choice of intervention and hypothesis and to predict major confounders and strategic design issues</td>
<td>Modelling: Identify the components of the intervention and the underlying mechanisms by which they will influence outcomes to provide evidence that you can predict how they relate to and interact with each other</td>
<td>Exploratory trial: Describe the constant and variable components of a replicable intervention and a feasible protocol for comparing the intervention with an appropriate alternative</td>
<td>Definitive trial: Compare a fully defined intervention with an appropriate alternative using a protocol that theoretically is defensible, reproducible, and adequately controlled in a study with appropriate statistical power</td>
<td>Long-term implementation: Determine whether others can reliably replicate the intervention in uncontrolled settings over the long term</td>
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<tr>
<td>USA: Sung et al. 2003, Crowley et al. 2004</td>
<td>Basic science research</td>
<td>Translation into humans: translational research and clinical trials</td>
<td>Health Services research and translation into clinical practice</td>
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<tr>
<td>USA: NIH Zerhouni 2003</td>
<td>Bench Basic science research: pre-clinical studies &amp; animal research</td>
<td>Translation to humans: phases 1 and 2 clinical trials</td>
<td>Bedside Human clinical research: phase 3 trials</td>
<td>Practice Clinical practice: delivery of care to right patient at right time</td>
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<tr>
<td>USA: The President’s Cancer Panel 2005</td>
<td>Basic science discovery</td>
<td>Early translation (1)</td>
<td>Late translation phase 3 trials</td>
<td>Adoption of advance by patients, providers and public</td>
<td>Payment mechanism(s) in place to enable adoption</td>
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<td>Promising molecule or gene target. Candidate protein biomarker. Basic epidemiological finding</td>
<td>Partnerships and collaboration (academia, government, industry) Intervention development Phase 1–2 trials</td>
<td>Regulatory approval Partnerships Production and commercialization</td>
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| UK: NIHR 2006 (National Institute for Health Research, 2006) | Translational medicine First in man to early clinical trials |

| USA: Westfall et al. 2007 | Bench Basic science research: pre-clinical studies & animal research | Translation to humans Phase 1 and 2 clinical trials | Bedside Human clinical research: phase 3 trials Practice-based research Phase 3 and 4 clinical trials, observational trials, survey research | Clinical practice |

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<tr>
<th>Thornicroft &amp; Tansella 2010</th>
<th>Basic science discovery</th>
<th>T1 Early human studies Early clinical trials</th>
<th>T2 Late clinical trials</th>
<th>Implementation</th>
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<tbody>
<tr>
<td></td>
<td></td>
<td>4.1 Adoption in principle</td>
<td>4.2 Early implementation</td>
<td>4.3 Persistence of implementation</td>
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MRC, Medical Research Council; NIH, National Institutes of Health.
Westfall translational step 2 refers to clinical guidelines, meta-analyses, systematic reviews.
Translational step 3 refers to dissemination research, implementation research.
Zerhouni translational step 2 refers to moving new medical discoveries into clinical practice.
(1) Dissemination: to community health providers and to patients and public.
these various formulations we propose a single overall schema, which consists of five phases (0–4) and three translational blocks (T1–T3), as shown in Fig. 1. This perspective integrates those elements cited in Table 1 and is proposed as a more coherent and comprehensive framework than other formulations to date.

**Phase 0: Basic science discovery**

Phase 0 refers to both basic laboratory and theoretical studies. The so-called ‘bench’ phase of basic laboratory research includes understanding therapeutic mechanisms of action, identifying promising molecule or gene targets and protein biomarkers, selecting candidate drugs and animal and laboratory (pre-clinical) studies. In terms of theoretical studies, this phase includes appraising relevant theories to ensure best choice of the candidate interventions, generating relevant hypotheses and anticipating the most important confounders, making judgements on the most critical research design issues, along with fundamental (aetiological) epidemiological research.

**Translational block T1**

The first translational block (T1) mediates the transfer of new understandings of disease mechanisms and drug actions gained in the laboratory into the development of new methods for diagnosis, therapy and prevention, alongside their initial testing in humans. In effect, the T1 block operates at the interface between animal and first-in-man studies.

**Phase 1: Early human studies**

In the drug development cycle, phase 1 studies are those that include healthy human volunteers and that aim to determine safety, tolerability, dose–effect relationships and early adverse effects. For psychosocial interventions, phase 1 refers to the period during which the key components of the intervention are identified, along with the manualization of the intervention.

**Phase 2: Early clinical trials**

For pharmacological interventions, phase 2 consists of early exploratory clinical studies to test efficacy in the target population; namely, individuals with the disorder to be treated. Studies of such patient volunteers can identify optimal doses, compare safety profiles with existing treatments and establish treatment duration. For psycho-social treatments, phase 2 investigations include exploratory studies (including randomized clinical trials), which describe the constant and variable components of a replicable intervention and which finalize a feasible protocol for comparing the intervention with an appropriate alternative (Campbell et al. 2007).

**Translational block T2**

In describing the US national clinical research enterprise, Sung et al. have distinguished the T1 and T2 translational blocks, where T2 refers to ‘the difficulty implementing therapeutic advances proven effective in large well conducted trials into the daily practice of medicine’ (Sung et al. 2003). In other words, T2 can be seen as the interface between efficacy and effectiveness trials, where the former are clinical studies carried out in ideal, experimental conditions, while the latter are those investigations conducted under routine clinical conditions (Tansella et al. 2006).

**Phase 3: Late clinical trials**

The next phase of clinical discovery refers to clinical studies of effectiveness and safety in target clinical populations (those with the condition to be treated), which are conducted over a longer time-scale and
which can identify less frequent and longer term side-effects (Tansella et al. 2006). In relation to ‘complex interventions’, phase 3 includes well-controlled investigations to compare a reproducible and fully defined intervention with an appropriate alternative under everyday clinical conditions, where the sample size is large enough to give a clear-cut answer to the primary question (Campbell et al. 2007).

**Translational block T3**

Westfall et al. have suggested a third gap (T3), at which evidence can fail to progress into clinical practice (Westfall et al. 2007). This is characterized as the distance between therapeutic interventions that are scientifically proven and applicable, for example, as formulated in clinical guidelines (Michie et al. 2007a), (phase 3), and the actual content of everyday clinical encounters (phase 4).

**Phase 4: Implementation**

The rapidly developing sector of ‘Implementation Science’ (Madon et al. 2007; Eccles et al. 2009) is beginning to identify the complex range of factors that interrupt the uptake of evidence-based practice at T3 in terms of: (i) the intention to implement; (ii) early implementation; (iii) persistence of implementation (Tansella & Thornicroft, 2009). Although this field is still at an early developmental stage, the journal Implementation Science is devoted to this field and has a growing scientific reputation. Nevertheless, although there are now thousands of published papers on the development of clinical guidelines across the range of healthcare, there are relatively few on how to put these guidelines into cost-effective, routine practice in any specialty (Institute of Medicine, 2001).

**Locating implementation science within translational medicine**

The overall purpose of translational medicine is ‘to test, in humans, novel therapeutic strategies developed through experimentation’ (Marincola, 2003). More specifically, translational medicine has been defined as ‘a discipline that increases the efficiency of determining the relevance of novel discoveries in the biological sciences to human disease and helps clinical researchers identify, through direct human observation, alternative hypotheses relevant to human disease. A further goal is to accelerate the rational transfer of new insights and knowledge into clinical practice for improving patients’ outcomes and public health’ (Littman et al. 2007).

The idea of translational medicine has been rapidly adopted in recent years and includes those studies that are related to: (i) defining the biology of disease; (ii) understanding the biological effects of therapeutics in humans; (iii) developing principles for the application of therapeutics to human disease; (iv) any clinical trial related to (i) – (iii) with an endpoint of toxicity and/or efficacy (Mankoff et al. 2004; Soderquest & Lord, 2010). From an historical point of view, the term translational medicine was until recently used in a somewhat broader sense, largely co-terminous with the whole range of the translational continuum described in this paper. It is only within the last decade that its use has been redefined more narrowly to refer to phase 1 (Marincola, 2003) or phase 1 and phase 2 (National Institute for Health Research, 2006) studies within the translational continuum.

To date, one common shortcoming of the conceptions of this whole translational pathway is that they are professionally driven, from left to right in Fig. 1. In other words, this vision is a simplified supply-side schema, in which scientists deliver inventions to clinicians (Perkins et al. 2007), who, in turn, deliver treatments to patients. Intriguingly, such thinking is not yet integrated with the conception of patient and public participation in healthcare. Specifically, at translational block T3, to date, there are few investigations about patient-related factors that accelerate or impede knowledge transfer. For example, well-informed patients are not only ‘stakeholders’ (for example, in developing clinical interventions or guidelines), but they also exert a powerful demand-side expectation for new treatments that are publicly understood to be beneficial, as has been clear in the HIV/AIDS field.

A related issue is the need to appreciate the distinction between dissemination and implementation (Rabin et al. 2008). The supply-side professional incentives that motivate scientists are primarily intended to disseminate their research findings via peer-reviewed journals, which are most often read by their scientific peers. There is a lack of clarity about who should have the responsibility and the resources to put such findings into clinical practice. In particular, there are few clear incentives for scientists to provide direct-to-patient information.

A further limitation of this field of study is that there is not as yet a clear overall theoretical paradigm for implementation science studies. Recently, however, there has been increasing attention to this theoretical deficit (Gardner et al. 2010; Michie et al. 2010; Webb et al. 2010), including a theoretically driven approach to understanding the formulation of clinical guidelines (Michie et al. 2007b). One integrative framework that has been recently proposed is the ‘Knowledge to Action model’, which considers three
states of knowledge (discovery, invention and innovation) and which is based upon the conceptual approach that stakeholders adopt and use knowledge that has perceived utility (Lane & Flagg, 2010). We anticipate a greater degree of integration in future implementation science studies between the theoretical approach used and the research designs employed (Craig et al., 2008).

In this paper, we have proposed a simple schema, consisting of five phases, to achieve a consistent terminology to understand the discovery, development, dissemination and implementation of new interventions. This schema also sets out three points at which communication blocks can occur. By better understanding the factors that promote or delay knowledge to flow across these blocks, we can accelerate progression along these translational medicine pathways and so realize earlier patient benefit.

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Declaration of Interest

None.

References


