Residual recurrence risk of ischaemic cerebrovascular events: concept, classification and implications

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INTRODUCTION
Patients with stroke still have substantial residual risk of stroke recurrence in spite of therapies following the guidelines.\(^1\) In addition to traditional vascular risk factors, conditions such as chronic inflammation, poor quality of care and variable response to standard therapy may contribute to the residual risk of stroke recurrence.\(^2\)\(^-\)\(^4\) Therefore, it is important to understand and classify the sources of residual risk to improve the prevention of stroke recurrence in patients already have a stroke.

Conceptualisation and classification of the residual risk
Residual risk is the amount of risk that remains after management or elimination of the known risk factors. Globally, the residual recurrence risk in patients who has a stroke has not been well estimated and varies between 3% and 6%.\(^5\)\(^-\)\(^6\) From the perspective of clinical practice, residual risk could be classified into three types:

Care-quality-based residual risk
Such type of residual risk developed mainly because of poor individual adherence to treatment. Though secondary prevention measures such as the use of antithrombotic, anti-hypertensive, anti-diabetic and lipid-lowering agents plus lifestyle modification have been recommended, adherence to these evidence-based therapies may be suboptimal in the real world.\(^7\) A multifaceted care quality intervention to improve adherence to guideline-based performance measures and secondary prevention measures among hospitalised patients can significantly reduce this type of residual risk.\(^7\) Another residual risk might be about achieving the target level of conditions such as in patients with hypercholesterolaemia. For example, low-density lipoprotein (LDL) cholesterol level was a common marker used to measure the effectiveness of cholesterol-lowering therapies. However, additional benefit from lipid-lowering agents beyond the recommended goals, such as an LDL level of 1.4 mmol/L or 0.78 mmol/L is not as clear, suggesting its role for residual risk.\(^8\)\(^-\)\(^9\)

Research-evidence-based residual risk
This type of residual risk refers to the conditions raised from bench research or only with limited evidenced from clinical research. For example, inflammatory conditions are known to be associated with stroke recurrence. The ongoing CONVINCE (Colchicine for Prevention of Vascular Inflammation in Non-cardio Embolic Stroke) study evaluates the effect of low-dose colchicine in reducing the rate of recurrent stroke in patients who had a stroke (URL: http://www.clinicaltrials.gov. NCT02898610). The result might shed some light. There are biological variations in the response to prevention therapies. For example, in those on clopidogrel, carriers of loss-of-function alleles of CYP2C19\(^10\)\(^-\)\(^13\) or protein biomarkers of glycated albumin,\(^14\) homocysteine\(^15\) and high-sensitivity C reactive protein\(^16\) may have a different response to clopidogrel. A recent study showed that residual risk was around 4.0% in a hypothetical scenario with further anti-inflammatory therapy and treatment for intracranial or extracranial arterial stenosis in patients who has a stroke under standard secondary prevention care (figure 1).\(^17\)

Intracranial atherosclerotic arterial stenosis is an important cause of recurrence stroke. There is limited evidence either stenting or aggressive medical therapy may prevent recurrent stroke in these patients.\(^7\) The benefit of revascularisation of occluded carotid artery still needs further evidence. In addition, psychological factors may correlate with recurrent stroke.
Unknown residual risk

Further research is needed to understand the unknown mechanism of recurrence and determine potential targets of intervention. Multiomics approaches may help with this effort by integrating data to better understand the relationship between different omics levels and their combined influence on underlying pathophysiology.\(^1\)

Implication and perspectives

Lowering the residual risk of stroke recurrence calls for comprehensive measures including improving the quality of care, facilitating the translation of research to clinical evidence and practice, and exploring new targets for prevention. First, a multifaceted intervention to facilitate care quality is needed to improve individual adherence to guideline recommended therapies. Second, facilitating the translation of evidence of research into clinical practice will help reducing the residual risk. Given the progressive decline in the rate of recurrence, it is likely that the large sample size is required.\(^1\) Appropriate design of the clinical trial and strong international collaborations would be required. Lastly, we must continue to explore new targets to manage the residual risks, which may be done through multiomics approaches (figure 2). Advances in technology have enabled the use of high-throughput techniques based on large screening processes that can minimise selection bias and generate extensive lists of molecules for targets of novel drugs.\(^1\) These high-throughput technologies, including genomics, transcriptomics, epigenomics, proteomics and metabolomics, enable omics studies to integrate millions of markers concurrently. Compared with a single layer of ‘omics’ that only provides limited insight into the biological mechanisms of stroke, multiomics approaches integrate data to better understand the relationship between different omics levels as well as their combined influence on pathophysiological processes.\(^1\)

Residual risk of stroke is a dynamic, evolving concept that may change around the recommendations of guidelines. However, standards of care are mediated through pathways not always directly addressed by our current management systems. Multiple new pharmacological agents targeting conceptually distinct pathophysiological targets have been explored to reduce residual risk. It is essential to continue this exploration of non-traditional risk factors through research, with a focus on multiomics approaches.

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Figure 1 Classifications of stroke residual risk.

Figure 2 Multiple omics for undetermined mechanism of stroke residual risk.
REFERENCES