Novel targets, treatments, and advanced models for intracerebral haemorrhage

Marietta Zille,* Tracy D. Farr, Richard F. Keep, Christine Römer, Guohua Xi, and Johannes Boltze*

aDepartment of Pharmaceutical Sciences, Division of Pharmacology and Toxicology, University of Vienna, UZA II, Althanstr. 14, Vienna 1090, Austria
bSchool of Life Sciences, Physiology, Pharmacology, and Neuroscience Division, Medical School, University of Nottingham, Nottingham NG7 2UH, UK
cDepartment of Neurosurgery, University of Michigan, Ann Arbor, MI 48109-2200, USA
dMax Delbrück Center for Molecular Medicine in the Helmholtz Association, The Berlin Institute for Medical Systems Biology, Berlin 13125, Germany
eSchool of Life Sciences, The University of Warwick, Gibbet Hill Campus, Coventry CV4 7AL, UK

Summary

Intracerebral haemorrhage (ICH) is the second most common type of stroke and a major cause of mortality and disability worldwide. Despite advances in surgical interventions and acute ICH management, there is currently no effective therapy to improve functional outcomes in patients. Recently, there has been tremendous progress uncovering new pathophysiological mechanisms underlying ICH that may pave the way for the development of therapeutic interventions. Here, we highlight emerging targets, but also existing gaps in preclinical animal modelling that prevent their exploitation. We particularly focus on (1) ICH aetiology, (2) the haematoma, (3) inflammation, and (4) post-ICH pathology. It is important to recognize that beyond neurons and the brain, other cell types and organs are crucially involved in ICH pathophysiology and successful interventions likely will need to address the entire organism. This review will spur the development of successful therapeutic interventions for ICH and advanced animal models that better reflect its aetiology and pathophysiology.

Introduction

Intracerebral haemorrhage (ICH) is caused by a loss of vascular integrity leading to bleeding within the brain parenchyma. ICH accounts for ~28% of all strokes and has the highest mortality rates among all stroke types. Hypertension is the most common risk factor of spontaneous ICH, exhibited by up to 70% of ICH patients and antihypertensive treatment significantly reduces ICH risk. Animal models may help to reveal underlying cellular and molecular mechanisms of vascular instability prior to first ICH occurrence. Mouse models of hypertension-related spontaneous ICH are available (8% high-salt diet + the nitric oxide synthase inhibitor L-NAME in drinking water on a double transgenic background of renin and angiotensinogen overexpression3; angiotensin II pump infusion + L-NAME in drinking water + injection of angiotensin II or norepinephrine in C57BL/6 mice).4 These models better reflect the more complex rupture of a blood vessel than what is modeled by cerebral collagenase injection.5 Major drawbacks are the relatively small ICH volume and thus comparatively short haemorrhage resolution time, as well as the unpredictable time of ICH onset.

Aetiology of ICH

Hypertension is the most common risk factor of spontaneous ICH, exhibited by up to 70% of ICH patients and

*Corresponding authors.
E-mail addresses: marietta.zille@univie.ac.at (M. Zille), johannes.boltze@warwick.ac.uk (J. Boltze).

www.thelancet.com Vol 76 Month February, 2022 1
cell degeneration alone may not be sufficient to induce blood vessel rupture. In mice with a mutation in the α1 chain of collagen type IV (COL4A1), a model of spontaneous ICH, hypermuscularization in the transitional segment between arterioles and capillaries led to an increase in intravascular pressure in the upstream arteriole that bursts at sites of smooth muscle cell loss. Similarly, combining angiotensin II-induced hypertension with low serum uric acid levels worsened the disruption of the smooth muscle cell-elastin contractile unit in cerebral vessels and ICH progression in mice. This highlights the importance of using spontaneous models of ICH for investigating aetiology.

Cerebral amyloid angiopathy (CAA) is characterised by amyloid β deposits in leptomeningeal and cerebral blood vessel walls decreasing vessel diameter and leading to microaneurysms which can cause ICH. Patients with CAA-related ICH also have a greater risk for ICH recurrence. CAA is mainly (80%) sporadic, but hereditary forms exist. Only the hereditary forms can be sufficiently modelled in animals (Swedish K670N/M671L and Dutch/Iowa E693Q/D694N mutations in the amyloid precursor protein), which to some extent can also mimic sporadic CAA. Whereas these transgenic mice display cerebral microbleeds, preclinical models of CAA leading to large ICH are lacking. Promising therapeutic approaches have been developed around amyloid β clearance.

Oral anticoagulant use increases the risk of ICH 7- to 10-fold. However, that risk is reduced in patients treated with direct oral anticoagulants (DOACs) compared to warfarin (Supplementary Table). Preclinical studies using collagenase- or laser-induced ICH in rodents have replicated the benefit of DOACs and the effectiveness of reversing anticoagulation was examined in animal models. In mice subjected to cerebral microbleeds, warfarin promoted deadly ICH, whereas DOACs increased microbleed burden without triggering long-term cognitive impairment. Whether anticoagulant treatment provokes ICH by aggravating existing microbleeds in humans remains unknown.

Vascular malformations including cerebral cavernous (CCMs) and arteriovenous malformations (AVMs) are major risk factors for ICH. CCMs are clusters of abnormal venous capillaries. There is no structural support of blood vessels from smooth muscle cells making CCMs vulnerable to rupture. Although most CCMs develop sporadically, several mutations causing CCMs have been identified (KRIT1/CCM1, malcavernin/CCM2, PDCD10/CCM3) and respective mouse models are available. AVMs are congenital entanglements of arterial vessels directly connected to the venous system without an intermediate capillary bed. Excessive vascular endothelial growth factor (VEGF) signalling is believed to promote AVMs, and Ras plays an important role for physiological VEGF signalling. Mutations in genes involved in the RAS/MAPK pathway such as KRAS have been observed in AVM endothelial cells in human patients. A mouse model of controllable Ras overactivation in endothelial cells has been created, in which cerebral AVMs and spontaneous ICH are observed.

Hypercholesterolaemia is associated with an increased risk for cardiovascular diseases, but decreases the risk for ICH in both sexes. However, hypercholesterinaemia can aggravate neuroinflammatory reactions after ICH similar to hypertension. An increased recruitment of neutrophils and monocytes is observed in dyslipidaemic mice, leading to poor functional outcome and exacerbated perihematomal oedema. Because of the high prevalence of hypercholesterinaemia in humans, further investigations in dyslipidaemic ICH models are warranted.

Haematoma expansion occurs in 20–40% of patients over the first day after ICH. Haematoma volume is a major determinant of outcome, and clinical trials have focused on limiting haematoma expansion. Phase III clinical trials (INTERACT2, NCT00716079; ATACH-II, NCT01176565; FAST, NCT00127283; FAST-EST, NCT03496883, Supplementary Table) have investigated pharmacological blood pressure lowering, including intensive approaches, after ICH (Supplementary Table). Current recommendations strongly suggest that elevated blood pressure should be treated as early as possible in patients with acute ICH.

A recent phase II trial (ICH-ADAPT II, NCT02281838, Supplementary Table) aims to identify the benefit of aggressive vs. conservative blood pressure lowering. It should be noted that studies focused on reducing blood pressure to limit secondary haematoma expansion after ICH have focused on reducing systolic or mean arterial blood pressure. It may be that fluctuations in pressure and blood flow play a role in inducing continued bleeding. Other trials to limit haematoma expansion include using haemostatic agents (tranexamic acid (TRANS-ACT, NCT03044184; STOP-MSU, NCT03389283) and recombinant Factor VIIa (FAST, NCT00127283; FAST-EST, NCT03496883, Supplementary Table)). Preclinical studies on haematoma expansion are hampered by available models. Yet, intracerebral injection of collagenase or autologous blood does produce a haematoma that initially expands and that can be exacerbated by hypertension and hyperglycaemia. However, there are concerns that the underlying mechanisms (e.g., gradual degradation of the endothelial basement membrane) do not reflect those occurring in ICH patients. A model using liquid polymer gel that coagulates on contact with tissue has recently been devised to create a mass that causes secondary bleeding, the extent of which is blood pressure-dependent. However, this model is lacking haemolysis-induced toxicity. Nevertheless, it may be useful for comparing different methods for limiting haematoma expansion.

The impact of physical haematoma evacuation (Figure 1) has been examined in many clinical trials
with, as yet, no evidence of improved neurological outcome (STICH II, ISRCTN22153967; MISTIE-III, NCT01827046, Supplementary Table). An alternate approach may be to accelerate endogenous haematoma resolution. Currently, in rodents, multiple agents such as peroxisome proliferator-activated receptor-γ and retinoid X receptor agonists have been shown to alter microglia/macrophage phenotype, enhance phagocytosis, speed haematoma clearance, and improve neurological outcome. Endogenous IL-4/Stat6 signalling is important in regulating haematoma resolution. Intranasal delivery of IL-4 nanoparticles also speeds resolution and improves neurological outcomes. An alternate approach is to block ‘don’t-eat-me’ signals expressed on erythrocytes that normally suppress phagocytosis (e.g., using a CD47 antibody). Whereas rodent ICH models have provided insight into mechanisms regulating endogenous haematoma clearance, studies on gyrencephalic species with larger brain sizes (e.g., pig, sheep) are needed to examine the effects of haematoma size and species physiological differences on endogenous clearance. Translation to the clinic may require a combination with physical evacuation to debulk the haematoma with accelerated endogenous clearance mechanisms to remove residual haematoma.

White matter (WM) injury with demyelination and axonal degeneration commonly occurs in human ICH. Despite this, WM injury has received less attention in preclinical ICH models in part because rodents have limited WM. Evidence indicates that ICH induces WM injury via multiple mechanisms including mechanical injury, oxidative stress (in part haemoglobin-/iron-mediated), neuroinflammation, excitotoxicity, and blood-brain barrier (BBB) disruption. Changes in the axonal cytoskeleton after ICH not only impact the physical structure of the axon but also mitochondrial transport and function leading to degeneration. Recent studies have suggested that the inhibition of histone deacetylases (HDACs) with scriptaid or conditional knockout of HDAC2 in microglia can reduce ICH-induced neuroinflammation and WM injury in mice. Other studies have targeted neuroinflammation with the antibiotic and inhibitor of microglial activation minocycline in piglets, and the sphingosine-1-phosphate receptor modulator FTY720/fingolimod in mice. Both approaches reduced WM injury. It should be noted that fingolimod is currently undergoing clinical testing for ICH (FITCH, NCT04088630, Supplementary Table). A pilot study on minocycline (MACH, NCT01805895, Supplementary Table) demonstrated that 400 mg of minocycline were safe and resulted in neuroprotective serum concentrations. Further clinical trials are needed to demonstrate the efficacy of minocycline in treatment of ICH.

Cell death is a hallmark of ICH. Whereas the haematoma causes immediate damage to cells, clot-derived

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**Figure 1.** The haematoma as a therapeutic target in ICH. The haematoma can cause brain injury via mass effect and the release of potentially toxic, plasma- or erythrocyte-derived factors. Surgical haematoma evacuation and accelerating endogenous haematoma clearance via phagocytosis are potential approaches, as is delaying erythrocyte lysis and inhibiting deleterious effects of clot-derived factors. RBC, red blood cell; MAC, membrane attack complex; Hb, haemoglobin; CA1, carbonic anhydrase 1; Prx2, peroxiredoxin-2; SIRPa, signal regulatory protein alpha.
breakdown products induce cell death hours to days after the initial bleed. Importantly, different cell death subroutines occur after ICH, including autophagy, necroptosis, and ferroptosis. Ferroptosis is an iron-dependent, non-apoptotic form of regulated cell death that is driven by lipid peroxidation and critically depends on glutathione peroxidase 4 (GPX4). Lipid peroxidation has been demonstrated in animal models of ICH. Knockout of 5-lipoxygenase improved functional recovery and N-acetylcysteine inhibited its toxic arachidonic products. Furthermore, the lipid peroxidation inhibitors ferrostatin-1 and liproxstatin-1 reduced neurological deficits, memory impairment, brain atrophy, lesion volume, and neuronal cell death in collagenase and autologous blood infusion models in mice. Increasing the expression of selenoproteins including GPX4 by selenium supplementation or a brain-penetrant selenopeptide abrogated ferroptosis and improved functional outcome in collagenase-induced ICH in mice. In ICH patients, ferroptotic gene expression is increased, but studies demonstrating functional benefit of anti-ferroptotic drugs for patients are still required. It should be noted that a recent analysis of the iDEF trial (NCT02175225, Supplementary Table) indicated a benefit of deferoxamine in patients with moderate (10–30 ml) haematomas.

Besides neurons, brain endothelial cells also undergo cell death after ICH. However, the underlying pathways are not yet fully elucidated. Endothelial dysfunction and BBB disruption have been demonstrated in collagenase and autologous blood infusion models of ICH in mice as well as in sheep in regions distant from the haematoma. The impairment of the BBB in ICH leads to brain oedema formation, the infiltration of immune cells, and the leakage of neurotoxic, pro-inflammatory, and vasoactive molecules.

ICH-induced cytotoxic and vasogenic brain oedema are intimately connected to parenchymal cell injury and vascular injury/BBB disruption. Currently, oedema treatments are generally limited to hyperosmotic solutions (mannitol/hypertonic saline) or hyperventilation. Both types of oedema are associated with a brain build-up of ions (e.g., Na⁺ and Cl⁻) and there has been considerable interest in the use of glibenclamide, an inhibitor of the sulfonylurea receptor 1-regulated ion channels, for reducing perihaeatomal oedema including a recently completed trial in ICH (GATE-ICH, NCT03741530, Supplementary Table). It should be noted that there is some disagreement in the preclinical literature about the effectiveness of glibenclamide in ICH.

Inflammation in ICH

Inflammatory events can be triggered by intraparenchymal blood (Figure 2), and partly resemble that in ischaemic stroke. They are, however, incompletely investigated and their temporal profile is not well understood. Local microglia and astrocytes are first...
responders and their proinflammatory activation promotes circulating immune cell influx, predominantly of macrophages. There is an increased release of inflammatory cytokines (e.g., interleukin-1β, tumor necrosis factor), free radicals, and chemokines. This attracts and activates lymphocytes. Importantly, these processes also contribute to perihaeamatomal oedema formation and potentially ICH growth by compromising BBB integrity. Most ICH models in immunocompetent animals allow investigating post-ICH inflammation impact and kinetics, but more specialised models exhibiting relevant comorbidities may provide better insights into inflammation post ICH.

Proinflammatory microglia/macrophages (often referred to as the M1 subtype) are considered promising therapeutic targets in early ICH as they drive neuroinflammation that is mainly associated with inferior outcome. Beneficial effects were observed with minocycline, a central nervous system (CNS)-penetrant tetracycline with inhibitory activity on pro-inflammatory microglia/macrophages, in rodent ICH models, but early-stage clinical studies were inconclusive. At later stages, anti-inflammatory (M2) microglia/macrophages subtypes exert beneficial effects for instance by contributing to haematoma removal as well as perihaeamatomal oedema resorption. M2 microglia/macrophages also promote local white matter integrity, and even exert neuroprotective effects. Thus, an alternative approach using human lactoferrin with the Fc domain of tetracycline with inhibitory activity on pro-inflammatory microglia/macrophages, in rodent ICH models, but early-stage clinical studies were inconclusive. At later stages, anti-inflammatory (M2) microglia/macrophages subtypes exert beneficial effects for instance by contributing to haematoma removal as well as perihaeamatomal oedema resorption. M2 microglia/macrophages also promote local white matter integrity, and even exert neuroprotective effects. Thus, an alternative approach using human lactoferrin with the Fc domain of minocycline, a central nervous system (CNS)-penetrant tetracycline with inhibitory activity on pro-inflammatory microglia/macrophages, may provide better insights into inflammation post ICH.

Inflammation and cytotoxicity driven by neutrophils are generally thought to exacerbate ICH injury, but neutrophils also exert beneficial and protective functions. For instance, the secretion of iron-scavenging lactoferrin can contribute to perihaeamatomal oedema reduction and increased haematoma clearance. Interleukin 27 reduces the amount of inflammatory/cytotoxic products in the neutrophil secretome and this increases the production of iron scavengers, thus shifting the balance in neutrophil action towards beneficial and protective effects. To note, this is a systemic process as the shift in the neutrophil secretome is induced in the bone marrow. Fusing human lactoferrin with the Fc domain of human IgG resulted in a molecule with a long plasma half-life and superior therapeutic outcome. To counterbalance inflammation, the injured CNS induces systemic immunosuppression. The overactivation of the sympathetic and parasympathetic nervous system in ICH results in spleen shrinkage and rapid lymphopenia. The degree of spleen shrinkage in ICH patients correlates with haematoma size, highlighting the link between ICH severity and immunosuppression, which in turn, predicts the likelihood of infections and impacts long-term outcome. Whereas the numbers of circulating (CD4+) T- and natural killer (NK) cells are decreased, there is a prominent invasion into the brain. Immunosuppression after ICH has been investigated in rodent ICH models using cerebral injection of autologous blood, revealing programmed death ligand 1 (PD-L1) and metoprolol as potential therapeutic options. However, haematoma location and size vary considerably in patients and may influence how the haematoma regulates immune responses after ICH. Hence, there is a need for future models resembling the clinical situation more closely. Of note, the immunosuppressive effects of fingolimod (FTY720), currently in clinical trial for ICH treatment (FITCH, NCT04088630, Supplementary Table), rely mainly on inhibiting helper (CD4+) and effector (CD8+) T cells as well as CD19+ B cells. Hence, post-ICH immunosuppression should be considered when designing immunomodulatory interventions for ICH.

Modulating immune cells in the CNS also affects the microbiome via the gut-brain-axis. The microbiome is well-studied in several CNS pathologies, including ischaemic stroke and plays a role in neuroinflammation, neuroplasticity and autoimmunity. However, little is known about the cross-talk between the gut and the brain in ICH. In collagenase-induced ICH in mice, reduced gastrointestinal motility and microbiota dysbiosis have been demonstrated, and adversely affected outcome. Recolonising ICH mice with healthy microbiota has shown promising results. It would be important to further investigate microbiome changes in ICH patients as well as microbiome transplantations. A human stool bank for healthy microbiota already exists (openbiome.org) and would be a powerful tool to leverage.

ICH sequelae and improving recovery
Cardiac complications are common after ICH and higher heart rate variability in the acute phase is associated with poorer 3-month outcomes. Preclinical studies using autologous blood or collagenase-induced ICH in mice have replicated cardiac complications and metoprolol reduced cardiac damage by abrogating sympathetic overactivation in addition to its immunosuppressive effects. Furthermore, splenectomy reduced cardiac dysfunction along with improving neurological outcome after autologous blood-induced ICH in mice. Due to their prevalence, cardiac complications should be assessed when testing novel therapeutic interventions preclinically.

Enhancing post-ICH recovery is an important aspect in therapeutic research. There is a plethora of well-
established tests of motor, sensory, and cognitive functions for rodent ICH models. However, evidence for cognitive deficits is limited and spontaneous recovery of motor function and compensatory mechanisms in rodents may overestimate the impact of therapeutic interventions after brain injury; some tests may not be able to discriminate between these effects. Sensitive tests such as automated gait assessment or kinematic measures can provide valuable insights into functional recovery. Interestingly, it seems that rehabilitation strategies predominantly focusing on gross motor function may not be optimal in rodents, an important parallel to human patients. However, more complex neurorehabilitation strategies are under-investigated for ICH. A combined application of enriched environment and task-specific motor training showed improved outcome in a rat model of striatal ICH more than a decade ago, but more detailed research on optimal neurorehabilitation strategies is currently missing. It is known that very early and intense rehabilitation can even impair functional recovery. This was shown in the phase III A Very Early Rehabilitation Trial after stroke (AVERT) which enrolled both patients with ischaemic and haemorrhagic stroke. Thus, timing and intensity of rehabilitation strategies requires further research. Interestingly, ICH location also impacts functional recovery. For instance, long-lasting deficits were observed after ICH in the internal capsule as compared to striatal lesions, despite a smaller lesion volume. ICH models targeting the internal capsule may therefore be well suited to investigate advanced rehabilitation or even restorative strategies. Detailed imaging protocols can be applied in ICH models and may reveal valuable morphological information related to functional recovery.

Depressive symptoms following ICH are common. Post-ICH depression may occur in up to 20% of patients and is associated with poorer long-term outcomes. Whereas treating depression is a priority, there is also an increased risk of secondary events associated with selective serotonin reuptake inhibitors. Furthermore, there has been interest in taking advantage of pleiotropic regenerative effects that drugs such as fluoxetine may offer. Three recent randomized clinical trials (FOCUS, ISRCTN8290762; AFFINITY, ACTRN1261000774921; EFFECTS, NCT0268213, Supplementary Table) assess fluoxetine for stroke recovery and all included ICH patients. They demonstrated that, whereas post-stroke depression was decreased, the risk of bone fractures and hyponatraemia was increased and functional outcome was not improved. It is clear that post-ICH depression represents an important unmet need and novel treatments may require improved knowledge of the underlying mechanisms. This effort is hampered by the sparsity of preclinical research. Some studies suggest rodents with ICH exhibit deficits in the elevated plus maze, sucrose preference, tail suspension, open field, and forced swim tests, but others have failed resulting in uncertainty whether animal models accurately reflect cognition and depression. Recent studies have investigated genetic links between ICH and depression and the results support the monoaminergic hypothesis, as well as the idea that pro-inflammatory cytokines may stimulate the hypothalamic pituitary adrenal axis.

Emerging therapeutic approaches

Small RNAs (size <200 nucleotides) are readily accessible in body fluids. The most abundantly investigated in ICH diagnosis and therapy are micro-RNAs (miRNAs, 20–25 nucleotides). Their expression profile not only distinguishes ICH patients from healthy controls but also ischaemic stroke and subarachnoid haemorrhage patients, which makes them valuable diagnostic (e.g., miR-124-3p) and prognostic markers (e.g., miR-130a). Dysregulated miRNAs and their putative mRNA targets are commonly associated with pathways regulating neuroinflammation, cell death, vascular smooth muscle and focal adhesion. Normalising imbalanced miRNA levels in ICH using miRNA mimics or antagonists leads to improved outcome after ICH in animals and in vitro models, including in models of the BBB. In addition, miRNA mimics and antagonists in ICH therapeutics commonly have anti-inflammatory functions, reduce perihematomatal oedema and haematoma size (miR130a, miR-223, miR-194-5p, miR-152, miR-7-5p), regulate BBB permeability (miR-130a, miR-27a, miR-126-3p), promote neuronal survival (miR-27a, miR-152), foster stem cell proliferation and migration, and improve endothelial function (miR-195). This suggests that miRNA-based therapies can be successful in improving ICH outcome in the clinical context when administered after ICH if appropriate delivery can be achieved.

Whereas the role and therapeutic potential of miRNAs is actively investigated, the potential of other small RNAs is relatively unexplored. These small RNAs include PIWI-interacting RNAs (piRNAs), transfer RNA derived small RNAs (tsRNAs), small nuclear RNAs (snRNAs), small nucleolar RNAs (snoRNAs), and small Cajal body RNAs ( scaRNAs). Seven tsRNAs have been demonstrated to be significantly changed in a rat model of collagenase-induced ICH and were involved in pathways participating in the oxidative stress response, endocytosis, and the regulation of G protein-coupled receptor signalling. Emerging clinical data highlights the differential expression of ribosomal and tRNA-derived fragments as well as snoRNAs in ICH compared with ischaemic stroke patients and healthy controls, respectively. The role of these dysregulated small RNAs in ICH is a field for future extensive studies and carries potential to open new avenues in understanding ICH pathobiology and developing ICH treatments.
Exosomes, also known as extracellular vesicles, are important mediators of regenerative mechanisms and exert therapeutic impact similar to that of regenerative cell populations. They are increasingly described in neurodegenerative disease, but knowledge in ICH is limited. Importantly, there is preliminary evidence for exosome-mediated anti-inflammatoryary mechanisms in mice and humans. Exosomes are frequently derived from mesenchymal stem cells (MSCs) as these cells are known to exert beneficial effects after ICH which are believed to be at least partly mediated by exosomes. Moreover, obtaining exosomes from MSCs is a well-established procedure. Specifically, exosomes obtained from bone marrow MSCs enriched with miR-146a-5p inhibited neuronal cell death and microglial M1 polarization compared to exosomes without enrichment. Exosomes from miR-19b-3p-mimic transfected adipose-derived stem cells were demonstrated to abrogate post-ICH ferroptosis in comparison to negative control mimic. Systemic delivery of miR-133b containing exosomes compared to miR-control reduced neurodegeneration by inhibiting RhoA and activating ERK1/2/CREB pathway even when administered 72 h after ICH, suggesting a clinical potential for this miRNA or exosomes as a delivery agent. A thorough characterisation of exosomes, including their cellular origin and content in the context of ICH will be important, and future research will also have to investigate therapeutic effects of exosomes derived from other (stem) cell populations.

Nanoparticle-based treatments, like many ICH-targeting therapies, have focused on mitigating pathogenic processes associated with the breakdown of blood in the brain. The most widely used approach has been to use nanoparticles to provide anti-oxidant benefits. Several nanoparticles have inherent anti-oxidant properties. For example, cerium oxide nanoparticles improved outcome in rodent models of ICH, with and without conjugation to polyethylene glycol (PEG), by reducing inflammatory activity and perihaematomal oedema formation. Several nanoparticle formulations are also amenable to functionalisation and/or modification to deliver therapeutic payloads. PEGylated hydrophilic carbon cluster nanoparticles have been bound to the iron chelator deferoxamine to further target ICH. Polymer-based nanoparticles remain popular as they are straightforward to manufacture, have multiple surface modification strategies, and are relatively stable. They have been modified to deliver oxidative therapies such as resveratrol and edaravone. The latter has potential for more rapid translation as edaravone has been used to treat ICH. Edavarone-containing nanoparticles further reduced perihaematomal oedema in patients with haematoma removal when compared to edaravone alone.

Limitations of animal models

As outlined above and summarized in Table 1, animal models have been used successfully to investigate some of the aetiologial and pathophysiological mechanisms of ICH, but all have limitations. There are indeed limitationsthat are common across animal models, such as an extraordinary spontaneous recovery of sensorimotor function in the rodent, as well as limited evidence for cognitive deficits. While translational validity is an important consideration, no model perfectly recapitulates the complex aetiology of spontaneous ICH in humans. Models should always be chosen with this in mind according to the specific research question. Furthermore, the inclusion of multiple comorbidities within models (including hypertension, dyslipidaemia, diabetes, arteritis) is recommended in order to better reflect the clinical scenario, and when testing new therapeutic approaches, the use of at least two models reduces considerations surrounding limitations and increases security that mechanisms of interest are widely applicable. Moreover, age and sex should be more often considered in animal studies as they impact the pathophysiology and clinical outcomes. In addition to behavioural/functional outcome assessments, non-invasive imaging methods such as magnetic resonance imaging may give further and longitudinal insights into the underlying mechanisms such as white matter injury, axonal degeneration, and haematoma expansion as well as when assessing potential treatment candidates. Thorough outcome analysis requires sufficient time to allow the lesion to be finally organized and for potential improvements to plateau. Post-injury/treatment surveillance time should be at least 3 weeks according to the Stroke Therapy Academic Industry Roundtable (STAIR) guidelines for preclinical research in ischaemic stroke using behavioural and structural or histological endpoints. Similar post-injury surveillance times are recommended for ICH.

Conclusion

Over the past two decades, it has become clear that ICH was previously neglected as a stroke type being distinct in its pathophysiology and treatment needs. We here summarized current therapeutical approaches and unaddressed targets that will help researchers in clinical translation (Figure 3). Importantly, there is increasing knowledge about the involvement of other cell types and organs that need more thorough investigation and to be considered for successful interventions. This also means that animal modelling needs to reflect this situation more closely to target the aetiology and pathophysiology of ICH better.

Outstanding questions

Whereas much has been learned about the pathological mechanisms involved in ICH-induced brain injury,
translating that information into novel therapies remains a challenge. Some of the mechanisms elucidated may have beneficial as well as detrimental effects (e.g., inflammation), the impact of which may vary with haematoma size and time after ictus. In addition, the importance of different injury mechanisms may differ between human and animal models. The use of large gyrencephalic species in

| Model | Aetiological or pathophysiological mechanisms addressed, translational value |
|-------|---------------------------------------------------------------------------|
| Autologous blood injection | - Endogenous haematoma clearance<sup>26,28,39,53</sup>  
- White matter injury and axonal degeneration<sup>10,200</sup>  
- Cell death<sup>30,31,32,103</sup>  
- Perihematoma oedema<sup>49,50,53,54</sup>  
- BBB impairment<sup>24,45,47,49</sup>  
- Inflammation<sup>28,45,51,101–103</sup>  
- Immunosuppression<sup>29</sup>  
- Cardiac complications<sup>51,54</sup>  
- Can be combined with comorbidities: angiotensin II infusion + hyperglycaemia (incl. haematoma expansion)<sup>22</sup>, hyperglycaemia<sup>105</sup> |
| Collagenase injection | - Anticoagulation<sup>22,53</sup>  
- Endogenous haematoma clearance<sup>26</sup>  
- White matter injury and axonal degeneration<sup>30</sup>  
- Cell death including ferroptosis<sup>37,38,40</sup>  
- Perihematoma oedema<sup>27,48</sup>  
- BBB impairment<sup>20,22,45,47,51</sup>  
- Inflammation<sup>20,51,52,103,104</sup>  
- Immunosuppression<sup>59</sup>  
- Gut-brain axis<sup>61</sup>  
- Cardiac complications<sup>59</sup>  
- Can be combined with comorbidities: dyslipidaemia, streptozotocin-induced diabetes<sup>20</sup>, spontaneously hypertensive animals (incl. haematoma expansion)<sup>5</sup> |
| Injection of blood components | - Axonal degeneration<sup>105</sup>  
- Cell death<sup>51</sup>  
- Inflammation<sup>51</sup>  
- White matter injury<sup>56</sup> |
| Laser-induced rupture of vessels | - Anticoagulation<sup>107</sup>  
- Microbleeds<sup>107</sup> |
| Cyclodextrin nanoparticle injection | - Anticoagulation<sup>14</sup>  
- Microbleeds<sup>14</sup> |
| Liquid polymer gel | - Hypertension (with additional phenylephrine injection)<sup>23</sup>  
- Anticoagulation (with additional anticoagulant injection)<sup>23</sup>  
- Mass effect, haematoma expansion<sup>23</sup> |
| High salt diet + L-NAME in drinking water in mice overexpressing human renin and angiotensinogen | - Reasonable modelling of the clinical situation  
- Spontaneous ICH  
- Hypertension as a risk factor for ICH<sup>3</sup> |
| Chronic angiotensin II infusion + L-NAME in drinking water + acute angiotensin II injection in mice | - Reasonable modelling of the clinical situation  
- Spontaneous ICH  
- Hypertension as a risk factor for ICH<sup>4</sup>  
- White matter alterations<sup>108</sup>  
- BBB impairment<sup>108</sup>  
- Inflammation<sup>108</sup>  
- Cognitive deficits and depression-like behaviour<sup>108</sup> |
| Chronic Angiotensin II infusion + L-NAME in drinking water + low serum uric acid levels in mice | - Reasonable modelling of the clinical situation  
- Spontaneous ICH  
- Hypertension as a risk factor for ICH<sup>7</sup>  
- Smooth muscle cell degeneration<sup>7</sup> |
| COL4A1 mutation in mice | - Reasonable modelling of the clinical situation  
- Spontaneous ICH  
- Hypermuscularization, smooth muscle cell degeneration<sup>7</sup>  
- BBB impairment<sup>109</sup> |
| CAA-related transgenic mice | - Reasonable modelling of the clinical situation  
- CAA<sup>9</sup>  
- Microbleeds<sup>9</sup> |
| CCM/AVM-related transgenic mouse models | - Reasonable modelling of the clinical situation  
- Vascular malformations<sup>5,14,18</sup> |

Table 1 (Continued)
addition to rodents for ICH modelling may address some concerns, but even they do not perfectly replicate human ICH size and time course. Thus, it is imperative to select the ICH model best suited to the respective research question or therapeutic target for best translational value.

Search strategy and selection criteria
Data for this review were identified by searches of MEDLINE, Current Contents, PubMed, and references from relevant articles using the search terms “intracerebral hemorrhage”, “hypertension”, “smooth muscle cells”, “hypercholesterolaemia”, “anticoagulants”, “haematoma expansion”, “haematoma evacuation”, “haematoma resolution”, “oedema”, “white matter injury”, “demyelination”, “axonal degeneration”, “ferroptosis”, “lipid peroxidation”, “brain endothelial cells”, “inflammation”, “small RNA”, “miRNA”, “exosomes”, “nanoparticles”, “cardiac complication”, “brain-heart axis”, “immunosuppression”, “gut-brain axis”, “recovery”, and “depression”. Only articles published in English were included with a particular focus on the past 3 years.

Contributions
All authors contributed to the writing of the article and approved the final version of the manuscript.

(A) Rodent models

| Model | Aetiological or pathophysiological mechanisms addressed, translational value |
|-------|--------------------------------------------------------------------------|
| Autologous blood injection in rabbits | Haematoma expansion, Haematoma evacuation, Perihematomal oedema, BBB impairment |
| Autologous blood injection in cats | Perihematomal oedema, Gyrencephalic brain, Higher white matter content |
| Autologous blood injection in dogs | Haematoma evacuation, Gyrencephalic brain, Targeted white matter injury possible, BBB impairment |
| Collagenase injection in dogs | Haematoma expansion and evolution, Gyrencephalic brain, Higher white matter content |
| Vessel puncture in dogs | Haematoma expansion, Cell death, Targeted lesion induction possible, Gyrencephalic brain, Higher white matter content |
| Autologous blood injection in pigs | Endogenous haematoma clearance, White matter injury, Cell death, Perihematomal oedema, Inflammation, Gyrencephalic brain, Higher white matter content |
| Autologous blood injection in sheep | White matter injury, Cell death, Perihematomal oedema, Gyrencephalic brain, Higher white matter content |
| Naturally occurring CAA in dogs | CAA, Gyrencephalic brain, Higher white matter content |
| CAA-related transgenic squirrel monkeys | Reasonable modelling of the clinical situation, CAA, Microbleeds |

(B) Non-rodent models

| Model | Aetiological or pathophysiological mechanisms addressed, translational value |
|-------|--------------------------------------------------------------------------|
| Autologous blood injection in rabbits | Haematoma expansion, Haematoma evacuation, Perihematomal oedema, BBB impairment |
| Autologous blood injection in cats | Perihematomal oedema, Gyrencephalic brain, Higher white matter content |
| Autologous blood injection in dogs | Haematoma evacuation, Gyrencephalic brain, Targeted white matter injury possible, BBB impairment |
| Collagenase injection in dogs | Haematoma expansion and evolution, Gyrencephalic brain, Higher white matter content |
| Vessel puncture in dogs | Haematoma expansion, Cell death, Targeted lesion induction possible, Gyrencephalic brain, Higher white matter content |
| Autologous blood injection in pigs | Endogenous haematoma clearance, White matter injury, Cell death, Perihematomal oedema, Inflammation, Gyrencephalic brain, Higher white matter content |
| Autologous blood injection in sheep | White matter injury, Cell death, Perihematomal oedema, Gyrencephalic brain, Higher white matter content |
| Naturally occurring CAA in dogs | CAA, Gyrencephalic brain, Higher white matter content |
| CAA-related transgenic squirrel monkeys | Reasonable modelling of the clinical situation, CAA, Microbleeds |

Table 1: ICH animal models to address aetiological and pathophysiological mechanisms and their limitations. AVM, arteriovenous malformations; BBB, blood-brain barrier; CAA, cerebral amyloid angiopathy; CCM, cerebral cavernous malformation; COL4A1, α1 chain of collagen type IV; ICH, intracerebral haemorrhage; L-NAME, N^\text{G}-nitro-L-arginine methyl ester.
Declaration of interests
The authors declare that they have no conflicts of interests.

Acknowledgements
We would like to thank Dr. Fan Xia the support in designing Figure 1 and Dr. Larisa Bulavina, a neuroscientist and professional medical illustrator (www.larisabulavina.com), for her invaluable help in graphical artwork.

Supplementary materials
Supplementary material associated with this article can be found in the online version at doi: 10.1016/j.ebiom.2022.103880.

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