Vascular disease and dementias: Paradigm shifts to drive research in new directions

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Abstract
Vascular disease was once considered the principal cause of aging-related dementia. More recently, however, research emphasis has shifted to studies of progressive neurodegenerative disease processes, such as those giving rise to neuritic plaques, neurofibrillary tangles, and Lewy bodies. Although these studies have led to critical insights and potential therapeutic strategies, interest in the role of systemic and cerebrovascular disease mechanisms waned and has received relatively less attention and research support. Recent studies suggest that vascular disease mechanisms play an important role in the risk for aging-related cognitive decline and disorders. Vascular disease frequently coexists with cognitive decline in aging individuals, shares many risk factors with dementias considered to be of the “Alzheimer type,” and is observed more frequently than expected in postmortem material from individuals manifesting “specific” disease stigmata, such as abundant plaques and tangles. Considerable difficulties have emerged in attempting to classify dementias as being related to vascular versus neurodegenerative causes, and several systems of criteria have been used. Despite multiple attempts, a lack of consensus remains regarding the optimal means of incorporating vascular disease into clinical diagnostic, neurocognitive, or neuropathologic classification schemes for dementias. We propose here an integrative, rather than a strictly taxonomic, approach to the study and elucidation of how vascular disease mechanisms contribute to the development of dementias. We argue that, instead of discriminating between, for example, “Alzheimer’s disease,” “vascular dementia,” and other diseases, there is a greater need to focus clinical and research efforts on elucidating specific pathophysiologic mechanisms that contribute to dementia phenotypes and neuropathologic outcomes. We outline a multitiered strategy, beginning with clinical and public health interventions that can be implemented immediately, enhancements to ongoing longitudinal studies to increase their informative value, and new initiatives to capitalize on recent advances in systems biology and network medicine. This strategy will require funding from multiple public and private sources to support collaborative and interdisciplinary research efforts to take full advantage of these opportunities and realize their societal benefits.

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1. Introduction

Improvements in public health and medical care during the 20th century led to substantial increases in life expectancy [1]. As a result, the principal causes of death have undergone a substantial shift from predominantly infectious
diseases to cardiovascular disease, cancers, and, increasingly, progressive neurodegenerative dementias [2]. With this increased longevity in industrialized societies, it has become clear that although some limited decline in certain cognitive functions with the aging process is almost universal, as many as half of all individuals living into their 80s and 90s undergo more severe cognitive and functional deterioration warranting a clinical syndrome diagnosis of dementia.

As the post–World War II “baby boom” generation now enters the vulnerable ages for dementia, increasing attention has been drawn to the medical and economic impact of dementias [3]. During this time, advances in scientific methodology and technological capabilities (brought about to a considerable extent by members of this same birth cohort) have greatly enhanced our ability to characterize the clinical and neuropathologic features of aging-associated cognitive disorders. In particular, the inventions of laboratory-based immunochemistry, molecular biology, and advanced microscopy as well as clinic-based magnetic resonance imaging, nuclear medicine imaging, and digital tomographic neuroimaging have revolutionized our understanding and diagnosis of brain diseases.

The principal disease processes leading to dementia in older adults can be broadly thought of as falling into 2 groups: (1) neurodegenerative processes occurring within brain tissue itself, and (2) vascular disease processes resulting in brain injury and dysfunction. The past half-century has seen dramatic growth in our understanding of neurodegenerative disease processes, most notably the processes underlying the formation of amyloid plaques and neurofibrillary tangles eponymously associated with Alzheimer’s disease (AD). In contrast, our view of the role of vascular disease processes in dementias has undergone several swings in emphasis over this interval, during which vascular mechanisms became less intensively studied. This shift in emphasis is evident even from the titles of some review articles on “vascular dementia” (VaD), alluding to this condition as an “enigma” [4] and asking whether we are on a “dead-end road” [5].

In this article, we will present a strategic overview of this extensive literature, highlighting some of the key findings that changed prevailing views regarding the role of vascular mechanisms in dementias. We will also consider the sources of research funding for studies conducted to date, and identify areas where gaps in funding priorities currently exist. We will then propose an agenda for future research in this area, including recommended modifications to the current paradigms by which vascular contributions to dementia syndromes are conceptualized.

As the historical aspects and evolution of concepts leading up to the present analysis have been well covered in several recent review articles and book chapters [6–9], we will only briefly summarize key points and will not recapitulate this material here, except where necessary to illustrate salient concepts.

2. Historical context

The notion that vascular factors mediate age-related cognitive decline was the prevailing view of “senile dementia” until the 1960s [4,10]. Thus, the terms “arteriosclerotic dementia” and its lay counterpart “hardening of the arteries in the brain” were in common use. Although this concept is often associated with the work of Binswanger first published in 1894 [6,11], many of the pathologic features can be traced back to the work of the French neurologist Max Durand-Fardel in the 1840s (cited in Ref. [12]). Until the last part of the 20th century, the occurrence of neuritic plaques and neurofibrillary tangles, the hallmark lesions described by Alzheimer in his seminal article of 1907 [13], were considered to be rare findings in patients with dementia, and more closely associated with early-onset or “presenile” dementias.

The emphasis in dementia research shifted dramatically starting in the 1970s and 1980s, based on several sets of observations. These include (1) the discovery that plaques and tangles were more common in postmortem material from individuals with dementing illnesses than had previously been thought, and more frequently than in those dying without dementia [14,15]; (2) the findings of neurodegeneration and associated deficits in the cholinergic system in the brains of patients with dementia [16], coupled soon thereafter with the discovery that anticholinesterase drugs had some therapeutic benefit in dementias [17] (refer also to subsequent meta-analysis in Ref. [18]); and (3) the discovery of the principal molecular composition of plaques (i.e., abnormally folded amyloid peptides [19]) and tangles (i.e., hyperphosphorylated tau proteins [20,21]). With the recognition that cholinergic-enhancing drugs have, at best, a limited impact on the underlying neurodegenerative processes [22], and the emergent preeminence of the amyloid cascade hypothesis [23], treatment development efforts have been intensely focused on interrupting the mechanisms of amyloid plaque formation. Several agents designed to affect amyloid production or clearance have been and are being tested in clinical trials [24]; however, none of these trials has yet progressed to the stage of applying for approval by Food and Drug Administration or other regulatory agencies.

3. Current state of knowledge

3.1. What is VaD?

VaD is often said to be the second most common form of dementia after AD [4,25]; some authors have even suggested it is the most common form [26]. However, such statements, and the data on which they are based, are predicated on having a commonly accepted definition of this condition. In the case of dementia attributable to vascular causes, this is far from the case. In fact, considerable variability exists in the literature with regard to the estimated prevalence of VaD, with estimates as low as 0.03% to as high as 98% (refer to Jellinger’s review in Ref. [4]). Most of the estimates range
between 11% and 18% for VaD, and between 22% and 34% for “mixed dementia,” which typically reflects features of both AD and vascular disease. Estimates are somewhat higher in series from Japan than in those from the United States and Europe.

This variability in the estimated frequency of VaD is due to at least three major factors: (1) there is no standard consensus definition of what constitutes VaD; (2) most studies have relied on clinical or autopsy samples, which are likely to be biased with regard to population estimates; and (3) an implicit assumption is made that VaD can be defined as a unique entity, existing apart from other conditions or diseases.

VaD is commonly understood as a condition in which cognitive impairment results from deleterious effects of vascular disease processes on brain structure and function. However, the kinds of vascular disease processes that can give rise to cognitive decline are highly varied (Table 2). These can include (1) cerebrovascular diseases (CVDs) involving either larger or smaller cerebral arteries, often (but not always) resulting in clinical stroke syndromes; and (2) cerebrovascular injury resulting from vascular events occurring outside the cerebral vasculature, such as emboli arising from the heart or large vessels proximal to the intracerebral circulation.

In contrast to AD, which has come to be defined as the progressive dementing syndrome associated with the accumulation of neuritic plaques and neurofibrillary tangles, it has been more difficult to define coherent syndromes of vascular disease processes leading to progressive cognitive decline and dementia. As a result, considerable debate has occurred not only about how to define VaD but also about what to call it. Thus, numerous terms have been used to describe dementia or cognitive impairment attributed to vascular disease involving the central nervous system (CNS; Table 1 in Ref. [4]).

### Table 1
Comparison of diagnostic criteria used for vascular dementia syndromes

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<tbody>
<tr>
<td>Stepwise deterioration</td>
<td>+</td>
<td>-</td>
<td>-</td>
<td>1</td>
<td>-</td>
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<tr>
<td>“Patchy” (unequal) distribution of cognitive deficits</td>
<td>+</td>
<td>-</td>
<td>+</td>
<td>-</td>
<td>4</td>
</tr>
<tr>
<td>Focal neurological signs</td>
<td>+</td>
<td>2</td>
<td>+</td>
<td>+</td>
<td>4</td>
</tr>
<tr>
<td>Focal neurological symptoms</td>
<td>+</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>4</td>
</tr>
<tr>
<td>Two or more ischemic strokes</td>
<td>-</td>
<td>2</td>
<td>-</td>
<td>3</td>
<td>-</td>
</tr>
<tr>
<td>Evidence of significant CVD</td>
<td>+</td>
<td>2</td>
<td>+</td>
<td>3</td>
<td>4</td>
</tr>
<tr>
<td>Etiologic relation to the disturbance</td>
<td>+</td>
<td>-</td>
<td>+</td>
<td>+</td>
<td>+</td>
</tr>
<tr>
<td>Temporal relationship between stroke and dementia</td>
<td>-</td>
<td>2</td>
<td>-</td>
<td>1</td>
<td>-</td>
</tr>
</tbody>
</table>

Abbreviations: DSM, Diagnostic and Statistical Manual for Psychiatric Disorders; ADDTC, Alzheimer’s Disease Diagnostic and Treatment Centers; ICD-10, International Classification of Diseases version 10; NINDS-AIREN, National Institute of Neurological Disorders and Stroke and European Association Internationale pour la Recherche et l’Enseignement en Neurosciences; CVD, cerebrovascular disease.

NOTE. Numbers in table refer to footnotes listed below.

1. either onset of dementia within 3 months after a recognized stroke and/or abrupt deterioration in cognitive functions, or fluctuating, stepwise progression of cognitive deficits; 2. evidence of >2 ischemic strokes by history, or neurological signs, and/or neuroimaging studies, or occurrence of a single stroke with a clearly documented temporal relationship with the onset of dementia and evidence of ≥1 infarct outside the cerebellum by CT or T1-weighted MRI; 3. either multiple strokes or a single strategically placed infarct; 4. focal neurological signs and symptoms or laboratory evidence indicative of CVD and judged to be etiologically related to the disturbance.

+ obligatory; -, not needed.


### 3.2. Clinical diagnostic criteria for dementias attributed to vascular diseases: Different criteria sets in use and limitations of existing systems

One of the first attempts to develop standard diagnostic criteria for VaD was included in the third edition of American Psychiatric Association’s Diagnostic and Statistical Manual for Psychiatric Disorders (DSM-III) [27], published in 1980. Three additional sets of criteria were published in the early 1990s. One of these was developed by the California Alzheimer’s Disease Diagnostic and Treatment Centers [28] and published in 1992; another was part of the International Classification of Diseases version 10 [29]; and the third was the result of a cooperative effort between the National Institute of Neurological Disorders and Stroke (NINDS) and the European Association Internationale

Table 2
Vascular diseases and conditions associated with cognitive impairment and decline*

<table>
<thead>
<tr>
<th>Poststroke dementia (heterogeneous pathobiology)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cortical infarct dementia (multi-infarct dementia)</td>
</tr>
<tr>
<td>Subcortical ischemic vascular dementia (Binswanger disease and lacunar state)</td>
</tr>
<tr>
<td>Strategic infarct dementia</td>
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<tr>
<td>Hypoperfusion dementia</td>
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<tr>
<td>Dementia caused by intracerebral hemorrhage</td>
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<tr>
<td>Dementia as a result of specific arteriopathies</td>
</tr>
<tr>
<td>Mixed dementia (vascular dementia and Alzheimer’s disease)</td>
</tr>
<tr>
<td>Vascular mild cognitive impairment (does not meet formal criteria for dementia)</td>
</tr>
</tbody>
</table>

behaviors and executive functions. Thus, the definitions of dementia included memory dysfunction as an obligatory feature. Dementias associated with vascular disease processes do not necessarily present with memory dysfunction as an entry criterion for VaD [36,37]. This is not entirely unexpected, given the substantial differences between criteria sets in terms of required elements, resulting in different subgroups of patients being captured. For example, the only element that is common to all these criteria sets is the evidence of CVD, although the definition of what constitutes CVD varies. Only DSM-III and NINDS-AIREN require a stepwise deterioration in function; only DSM-III and International Classification of Diseases version 10 require a “patchy” distribution of cognitive deficits; and the criteria vary in their requirement of a temporal relationship between cerebrovascular events and cognitive decline. In addition, clinical judgment is required by many of the criteria, for example, of an “etiologic” relationship between cerebrovascular events and cognitive impairment.

3.2.1. Biases in definitions of “dementia” and “cognitive impairment”

One important difficulty with the study of vascular mechanisms in dementia is that until relatively recently, the definitions of dementia included memory dysfunction as an obligatory feature. Dementias associated with vascular disease processes do not necessarily present with memory dysfunction, often manifesting initially by alterations in behaviors and executive functions [38]. Thus, the definitions of dementia and “mild cognitive impairment” (MCI) needed to be broadened to encompass syndromes in which memory dysfunction is not prominent, but other cognitive domains are substantively impaired. Subsyndromal distinctions within this broader category (MCI) can thus be made, depending on the specific cognitive domain(s) affected. Importantly, assessment batteries used in neurocognitive evaluation of dementia need to be modified to include more detailed assessment of the domains of executive function, such as attention, working memory, planning, and set shifting, as well as speed of processing. Standard neurocognitive testing protocols of varying lengths ranging from 5 to 60 minutes have been proposed [39].

The notion of MCI involving domains exclusive of memory was also recognized by subdividing MCI into amnestic and nonamnestic subtypes [40]. Recent studies suggest that vascular disease processes are important contributors to nonamnestic MCI phenotypes [41,42]. In addition, the DSM-V Work Group has proposed criteria for cognitive impairment that will not require memory dysfunction [32]. Nevertheless, the bias toward memory impairment as an “entry criterion” for VaD or cognitive impairment in previous diagnostic schemes must still be taken into account in the evaluation of studies conducted using those criteria.

Finally, there are sociocultural and nosologic trends within medicine, arguing for a shift in terminology and classifications of dementias. As the technical term dementia has gained wide public usage, it has taken on pejorative and potentially stigmatizing connotations [43]. This has led to the current moves to eliminate the term from the medical lexicon. Some authors have adopted the term “vascular cognitive impairment” [7,44]; however, this is variously used to refer either to frank dementia syndromes or all cognitive impairment attributed to vascular causes [45]. Although still controversial, the DSM-V Work Group has adopted the term “major neurocognitive disorder” for dementia and “minor neurocognitive disorder” for MCI, which are then subclassified according to suspected etiology (e.g., AD, vascular neurocognitive disorder, Lewy body disease, “mixed”). Increasingly, the term vascular cognitive impairment is being used, with the notion that this encompasses both dementia and MCI due to vascular brain injury [7,39,45–48]. Such efforts are critical to improve specificity and acceptable use of diagnosis as knowledge evolves but do present challenges for comparative interpretation.

3.3. Neuroimaging findings: Relationship with vascular and brain lesions

As neuroimaging methodologies became more widely available and were applied to dementia research, patients with dementia and CVD were often found to have areas of altered appearance in white matter areas. These were initially observed on X-ray computed tomography as areas of hypodensity, leading to the introduction of the term “leukoaraiosis” by Hachinski et al [49]. As the use of magnetic resonance imaging became more widespread, areas of increased signal intensity on T2-weighted scans, now termed “white matter hyperintensities” (WMH), were observed in similar patterns. Subsequently, these areas were more clearly shown in some cases using sequences such as fluid-attenuated inversion recovery (to attenuate the cerebrospinal fluid signal and thus better delineate periventricular hyperintensities) and proton density imaging.

These white matter alterations are typically considered in terms of their location, that is, a distinction is made between those observed in periventricular regions and those observed in other subcortical regions [50]. The number and size of these lesions were found to increase with age [51] and were more prevalent in those with dementia than those without; among those with dementia, WMH were more frequent...
in those diagnosed with vascular than Alzheimer-type dementias (AD) [52].

Although WMH show a strong association with CVD [53], they are also increased in those with mood disorders, such as bipolar disorder [54], and migraine [55]. Whether these indicate CVD and, if so, what type of CVD and injury, is not known, as WMH due to different underlying mechanisms may be indistinguishable using standard neuroimaging approaches. Such findings may complicate the interpretation of WMH observed in patients with co-occurring conditions.

3.3.1. Lack of standardization of neuropathologic criteria for vascular disease in brain

A major handicap to diagnosis and research on VaD is the lack of a widely accepted, standardized, neuropathological approach to the identification, categorization, localization, and, in particular, quantitation of CVD and vascular brain injury. Common approaches used by major research groups have consisted of characterizing CVD according to the presence or absence of large- or small-vessel infarcts and lacunae evident on gross inspection of coronal sections, the presence and severity of cerebral amyloid angiopathy (CAA), and the presence and severity of hippocampal sclerosis. Less consistently applied have been the notations of atheromatous disease in the circle of Willis, microinfarcts, microhemorrhages, perivascular expansions, lipohyalinosis, microaneurysms, laminar necrosis, or granular atrophy. Indeed, the National Alzheimer’s Coordinating Center Neuropathology Diagnosis Guidebook [56] includes in its assessment items “meant to indicate the presence of vascular pathology, but not the absolute burden, volume, or severity of change” (p. 5).

Of particular importance for current concepts of VaD is the absence of any standardized neuropathological approach for classifying and measuring deep white matter and periventricular disease and injury. Furthermore, there have been no widely used systematic classification schemata that incorporate lesion location. Clearly, a small volume of vascular infarction in the medial thalamus will likely have very different and potentially more significant effects on cognitive function than similar or even much larger injuries in posterior white matter.

Arterial CVD encompasses a broad range of pathological processes (e.g., atheromas, hyalinosis, arteritides, aneurysm) that occur in large extracranial and intracranial blood vessels, small arteries and arterioles, and microvasculature, including capillaries. Cerebral thrombosis, embolism, and hemorrhage at any of these levels of the cerebral arterial vasculature are the principal proximal causes of ischemic brain injury. In addition, an underinvestigated pathology in the realm of aging that is unique to the brain is the breakdown of the blood–brain barrier. This may be particularly relevant to the promotion of neurodegeneration independently, additively, or synergistically with AD and other common neurodegenerative disease processes.

3.4. High rate of co-occurrence of Alzheimer-like and vascular disease–related neuropathologic features on postmortem examination of individuals with dementia

CVD is common in the population; its risk increases with advancing age, as does that for AD and other neurodegenerative dementias. Neuropathologic evidence of chronic vascular disease can take various forms, reflecting the diverse ways in which the brain can be affected by vascular disease processes. These include large-vessel infarcts with regional or lobar encephalomalacia, small-vessel macroscopic cystic/lacunar infarcts (typically <1 cm in white matter or deep cerebral nuclei), microinfarcts in gray or white matter (observable only microscopically), leukoencephalopathy (microscopically observable rarefaction of white matter), cribriform changes (perivascular space with ischemic gliosis), and CAA.

Although physicians and scientists would prefer to apply a singular etiologic diagnosis for dementia, it is readily apparent that multiple pathologies in the brain are the rule rather than the exception. Large-scale epidemiologic clinicopathologic programs around the world have convincingly demonstrated that many, or even the majority of, people with dementia have mixed pathologies, chiefly AD with infarcts and, to a lesser degree, Lewy bodies [57–67]. However, it is difficult to glean from the extant literature just how common pathological concurrence is, given the widely varying methods for identifying, classifying, and reporting AD and vascular disease frequencies in the same patient populations.

The frequent co-occurrence of “Alzheimer-like” features and stigmata of vascular disease can make it difficult or impossible to attribute cognitive decline to one process or the other. However, if headway can be made in determining the degree to which vascular pathology contributes to cognitive impairment, systematic accounting of this pathology along the dimensions of neuropathological type, size, location, and perhaps chronicity will need to be incorporated into standardized postmortem assessment.

Although postmortem examination of the brain is the most definitive means of identifying structural changes in dementias, and has provided invaluable information regarding these changes, its intrinsic limitations must also be kept in mind. Postmortem examination is inherently a static, post hoc, and correlative measure that necessarily reflects the end result of pathologic processes that typically have progressed over many years. In contrast, we know that the brain shows considerable plasticity throughout adult life [68,69], and thus has the ability to compensate for the effects of a wide range of disease processes [70]. As a result, pathologic examination alone does not provide information regarding the degree to which the abnormalities observed are responsible for the clinical features observed during life. Thus, neuropathologic evidence of vascular disease may not have been associated with clinical evidence of cognitive impairment; conversely, cognitive impairments may be observed during life that do not have a clear counterpart on postmortem brain examination [71]. This difficulty in extrapolating between
clinical dysfunction and neuropathologic stigmata is reflected in the recent proposed revision of diagnostic criteria for AD [72], in which clinical and neuropathologic diagnoses are made independently. Longitudinal studies including both in-life measures and postmortem evaluation are critical to better define these relationships. Furthermore, both clinical and neuropathologic criteria for vascular disease need to be standardized by a consensus process so that results will be comparable across institutions and over time.

3.5. Overlap in risk factors for “vascular” and “Alzheimer-type” dementias

We are increasingly recognizing that risk for AD is increased by the factors traditionally associated with cardiovascular diseases. Even the major single gene associated with increased AD risk, namely, that for apolipoprotein E (APOE), was earlier identified as a risk gene for cardiovascular diseases [73]. The modifiable risk factors for which the best evidence exists include obesity during midlife, type 2 diabetes (T2DM), and cigarette smoking [74] (Table 3). Of these, the co-occurrence of obesity (in particular, visceral or intra-abdominal obesity) together with insulin resistance or glucose intolerance, subclinical degrees of elevated blood pressure, and elevated total or low-density lipoprotein (LDL) cholesterol (LDL-C) is associated with a comparable increase in risk for adverse cardiovascular outcomes as a threshold-level degree of any one of these factors [75] and has been called the “metabolic syndrome.”

Intensive study of this syndrome over the past 10 to 15 years has shown that these various components mutually interact with each other and with other factors such as inflammatory responses, alterations in vascular endothelial function, platelet activation, and engagement of hemostatic and antithrombotic mechanisms to contribute to the progression of atherosclerotic and atherothrombotic disease, leading to adverse cardiac outcomes such as myocardial infarction and death [75]. Although there is ongoing debate as to the core components of this syndrome, the precise mechanisms involved, and the implications of the overall syndrome for cardiovascular risk apart from those of its constituent risk factors [76–78], there is general agreement that these risk factors are frequently associated and can exert mutually reinforcing influences on each other.

These observations draw attention to the potential role of vascular disease mechanisms in AD, traditionally conceptualized as a dementia characterized exclusively by the neuropathologic stigmata of plaques and tangles. A major question is whether dementias related to vascular disease processes and those related to amyloid- and tau-related processes are fundamentally different, producing additive effects on cognitive function, or whether the co-occurrence of both disease processes has synergistic effects on cognitive failure. Such questions require careful longitudinal assessment of both vascular and neurodegenerative disease measures in conjunction with a broad sampling of neurocognitive functions, imaging and biochemical biomarkers, and measures of functional status, with postmortem assessment of both the cerebral vasculature and brain parenchyma for correlation with in-life measures. Animal and cell-based models are also needed to test specific mechanistic hypotheses of interactive effects.

4. Paradigm shift for future research: Focus on risk factors, multiple and interacting mechanisms, and disease modification

It should be clear from the above discussion that (1) there is considerable overlap in the risk factors for what is now termed “Alzheimer’s disease” and VaD or vascular cognitive impairment; (2) there is a lack of consensus regarding clinical criteria for cognitive impairment related to vascular disease, including even what to call it; and (3) postmortem examination of individuals with dementia commonly shows features of both AD (i.e., plaques and tangles) and VaD (i.e., CVD). We propose here that one major obstacle that impedes future research is the focus on defining and distinguishing these entities as discrete and independent diseases. We may not precisely understand the relative contributions each “lesion” (however it is defined) makes to cognitive decline, but we can appreciate how much they overlap, and thus presume they are intimately interactive. As a result, we propose an alternative approach of distinguishing between risk factors, pathophysiologic processes, clinical syndromes, and neuropathologic findings observed at autopsy. We believe that such an approach permits the development of a useful blueprint for future research on the causes, treatment, and, ultimately, prevention of these diseases.

4.1. Risk factors—General comments

Risk factors are defined as characteristics of individuals that are associated with statistically valid predictions about
the likelihood of a specific outcome (usually a disease or event such as cancer, stroke, myocardial infarction, or death) at some future time [79]. The complementary, albeit less frequently used, term “protective factor” denotes characteristics associated with a reduced likelihood of a future outcome.

It is important to bear in mind that risk and protective factors, being largely identified through observational studies, are correlational and do not directly identify a mechanism of disease; however, they can provide important clues about underlying mechanisms that can be tested by interventional studies designed to correct the risk factor.

As indicated in the Introduction section, CVDs have long been implicated as risk factors for dementias, and were considered their primary causes at one time. CVDs are now considered as part of the broader framework of atherothrombotic vascular disease (ATVD), which also includes coronary artery disease (CAD) and peripheral vascular disease. In turn, the most common modifiable risk factors for ATVD include systemic hypertension, T2DM, and dyslipidemia [80]. We now discuss these specific conditions with regard to their potential roles as risk factors for dementia and cognitive decline.

4.2. Cardiovascular risk factors

4.2.1. Stroke and CVD

The occurrence of a stroke is associated with a doubling of the risk of developing dementia [81]. A single stroke can cause dementia if it causes sufficient damage to brain areas or pathways important for cognitive functions (refer to review in Ref. [81]); risk estimates vary across studies and are difficult to compare because of different patient sampling methodologies, follow-up intervals, and criteria used for diagnosis of dementia. The risk of dementia appears to increase with recurrent strokes and with the volume of brain tissue sustaining lasting injury, although there is considerable heterogeneity within this general framework. Small strokes in areas critical for cognitive functions can result in greater impairment than expected; this is often termed “strategic infarct dementia.” Most of the literature on dementia associated with “strategic” strokes is published in the form of single case reports or small case series (refer to Ref. [81] for review).

The development of cognitive decline in association with the occurrence of multiple small strokes or lacunar infarcts was termed “multi-infarct dementia” by Hachinski et al [82]. Typically, patients manifesting these features have chronic hypertension and/or diabetes, which results in damage to smaller arteries in the cerebrovascular system and associated damage to subcortical brain structures and white matter tracts. These lacunar infarcts appear to correspond closely to white matter abnormality intensities observed with contemporary neuroimaging methods [49].

4.2.2. “Silent strokes” and microbleeds

Postmortem examination of the brain frequently reveals evidence of small infarcts [83] and hemorrhages [84] that can occur in the absence of macroscopic strokes and that were not clearly associated with acute or chronic stroke symptoms during life. Although neuroimaging methods are becoming increasingly sensitive to such lesions, many remain beyond the resolution of current technology [85]. There is some evidence that these lesions may be associated with cognitive impairment [83], although the nature and extent of their direct contribution to cognitive impairment have not been elucidated.

4.2.3. Hypertension

Systemic hypertension is one of the best-known risk factors for CVD and stroke [86]. The risk of stroke increases by approximately 30% for each 10-mm Hg increase in systolic blood pressure >115 mm Hg, and for each 5-mm Hg increase in diastolic blood pressure >75 mm Hg; a concomitant decrease in stroke risk is observed for lowering of blood pressure with treatment [86]. Hypertension, especially during midlife, has also been shown to be an important risk factor for cognitive impairment and dementia [87]. The etiology of most forms of hypertension is not known with certainty. Family history and racial/ethnic background are known nonmodifiable risk factors. Some single-gene variants are beginning to be identified based on genome-wide association studies, although the effects of any given risk allele tend to be small [88]. Modifiable risk factors include salt intake in some cases and other lifestyle factors such as physical activity. In many cases, dietary and lifestyle modifications are not sufficiently effective in lowering blood pressure, and medications must be used. Numerous classes of medications are now available to treat hypertension, and algorithms have been developed to guide the use of these medications [89].

4.3. Metabolic risk factors

4.3.1. Type 2 diabetes mellitus

T2DM, an important risk factor for CVD, is associated with a 1.5- to 2-fold increase in the risk of dementia [90], and this is independent of stroke [90]. The incidence of T2DM is increasing in developed countries [91]. This trend has been attributed in part to the dramatic increase in the number of overweight and obese individuals, as weight gain is a key risk factor for the development of T2DM. The diagnosis of T2DM is based on fasting blood glucose levels of >126 mg/dL; individuals with fasting blood glucose >110 mg/dL are considered to be “at risk.” However, it is recognized that elevated fasting glucose most often results from increasing degrees of insulin resistance developing over many years, leading to alterations in glucose homeostasis and energy metabolism, and culminating in a failure of pancreatic insulin secretion to maintain blood
glucose within the normal range. Accordingly, the development of fasting hyperglycemia is preceded by elevated postprandial blood glucose or glucose intolerance. The progression of glucose intolerance to fasting hyperglycemia can be prevented in some cases by medications or, perhaps even more effectively, by lifestyle interventions [92].

Diabetes is associated with pathologic changes in both large (macrovascular) and small (microvascular) blood vessels. Some of the pathologic consequences of T2DM are attributed to the effects of sustained hyperglycemia, whereas others are related to the preceding phase of hyperinsulinemia and its adverse effects. Still other morbidities have been proposed to be related to the episodes of hypoglycemia that can occur as a consequence of treating T2DM or its prodromal phases with hypoglycemic agents and/or insulin.

4.3.2. Metabolic syndrome: Effects of interactions among multiple risk factors

Epidemiologic data had suggested specific thresholds for the effects of the major risk factors on CAD risk [75] (e.g., total cholesterol and LDL-C, blood pressure, fasting and postprandial glucose). In the course of analyzing these data, it became apparent that subthreshold levels of multiple risk factors jointly conferred levels of risk similar to those from threshold levels of any one risk factor. These findings led to the concept of the “metabolic syndrome,” [75] in which multiple pathophysiologic mechanisms interact to produce adverse outcomes. This concept continues to evolve [76–78], and its components and relevance to the development of cognitive impairment owing to vascular disease will be discussed further later in the text.

4.4. Vascular pathophysiologic processes contributing to dementia and cognitive decline

The extensive research on mechanisms underlying the development and progression of atherosclerotic and ATVDs offers a number of potential mechanisms likely to contribute to the impact of such diseases on cognitive function during the aging process. We highlight here some of the key findings from this research. In doing so, we recognize that many of these processes exert mutual influences on one another, whereas other processes may be involved that are not delineated here; thus, treating them separately may understate the degree of interactive and dynamic effects that may be operating in vivo. It should also be noted that this is a rapidly advancing field, and it is critical for future research on vascular mechanisms in dementia to integrate and apply these findings to inform our understanding and to identify and test new targets for prevention and treatment.

4.4.1. Aging-associated biomechanical changes

Aging is associated with increased stiffness of arteries as a result of fragmentation and depletion of elastin and increased deposition of collagen in the vessel wall [93,94]. These changes primarily affect larger arteries and reduce the ability of these arteries to respond to regulatory influences. In addition, the increased stiffness reduces the ability of larger arteries to buffer the pulse wave that occurs with cardiac contraction, and can thus result in greater mechanical strain on smaller arteries. Finally, many of the risk factors for vascular disease, such as hypertension, widened pulse pressure, diabetes and metabolic syndrome, and inflammatory processes and oxidative stress associated with atherosclerosis, can exacerbate the effects of aging on vascular stiffness [95].

4.4.2. Atherosclerotic plaque formation and progression

Atherosclerosis is a disease process that develops over many years, beginning with the formation of fatty streaks in large arteries during adolescence [96]. This process can be conceptualized as occurring in stages. The initial stages in atheroma formation involve uptake of LDL particles into the intimal layer of arteries. These LDL particles become modified by oxidation and glycation, inducing local inflammatory responses. The inflammatory foci attract macrophages through secretion of chemokines and chemoattractant factors, leading to a cyclic process of oxidative stress and release of inflammatory mediators. The next major step in this process involves entry of smooth-muscle cells into the intima, enlarging the plaque and leading to fibrosis and the formation of a fibrous cap. This plaque can be stable for extended periods, until an event occurs to cause rupture of the plaque, releasing its contents into the circulation and resulting in the occlusion of one or more downstream blood vessels. In the cerebral vasculature, this will precipitate a stroke or other cerebrovascular event if the occluded blood vessels are sufficiently large to cause compromise of brain function. Vascular occlusions involving smaller territories or “silent” areas of brain may not cause acute symptoms.

4.4.3. Insulin resistance and related components of the metabolic syndrome

As indicated earlier in the text, the occurrence of subthreshold levels of several risk factors for atherosclerotic cardiovascular disease (chiefly CAD) had been found to confer a degree of risk equivalent to that of any one risk factor present at threshold levels [75]. The key elements of risk identified in this analysis were (1) excessive visceral adiposity, as assessed by measuring waist circumference; (2) an atherogenic pattern of alterations in serum lipid measures (i.e., elevated total and LDL-C, elevated triglycerides, and decreased high-density lipoprotein [HDL] cholesterol [HDL-C]); (3) elevated blood pressure; (4) insulin resistance, with or without glucose intolerance; and (5) evidence of a prothrombotic and proinflammatory state.

The pathophysiologic basis underlying the associations between these factors is an area of active investigation and continued controversy [76–78,97]. The most prevalent view is that visceral obesity leads to systemic insulin resistance, which in turn results in a number of deleterious consequences that promote atherogenesis and progression
of atherosclerotic lesions, and eventually increases the risk of acute ischemic events due to compromise of blood flow and oxygenation to susceptible tissues such as the heart and brain [98,99].

4.4.4. Relationship between (visceral) obesity and insulin resistance

Obesity, particularly that involving intra-abdominal regions, is associated with an increased risk of insulin resistance [100,101]. In large studies of apparently healthy individuals, body mass accounts for approximately 25% of the variability in insulin sensitivity [102]. A number of potential mechanisms have been implicated in this effect. These include alterations in lipid metabolism [103], increased production of proinflammatory cytokines [104], and increased arterial blood pressure [105].

Some authors have questioned whether the driving force for insulin resistance is visceral obesity or whether obesity in general is sufficient [77]. Thus, there is evidence that intra-abdominal adipose tissue can be regulated differently than that in subcutaneous regions [106], shows differential expression of proinflammatory cytokines [106] and chemotactic factors [107,108], and in obese subjects shows increased infiltration of macrophages and other inflammatory cells [109,110]. Free fatty acids released by lipolysis in visceral adipose tissue are delivered directly to the liver via the hepatic portal system [111], which can inappropriately increase hepatic glucose production [112] and may contribute to nonalcoholic hepatic steatosis [113]. However, in a recent study, surgical removal of omental fat did not improve insulin resistance or other metabolic parameters [114]. Further research is needed to better define the role of different fat depots in the mechanism of insulin resistance associated with weight gain. Nevertheless, there is strong reason to believe that the dramatic global increase in the number of overweight and obese individuals over the past several decades will substantially increase the long-term burden of cognitive impairment in addition to the metabolic and vascular diseases attributable to insulin resistance. Further understanding of the underlying mechanisms will aid in identifying the individuals most at risk for such consequences and more effective targeting of interventions to mitigate these risks.

4.4.5. Intrinsic brain insulin resistance in AD

There is also emerging evidence that neuronal insulin resistance may be an important feature of AD, independent of systemic T2DM [115]. A number of postmortem and clinical biomarker studies have now documented abnormalities in insulin and insulin receptor levels in the brain as well as major changes in intraneuronal insulin signaling pathways that together demonstrate a state of intrinsic insulin resistance in the brain in AD. Some have even proposed that this may be called a “type 3” diabetes mellitus, although it is not clear whether deficient neuronal insulin signaling is directly associated with alterations in glucose metabolism as would be implied by this designation [116].

4.4.6. Alterations in circulating lipids and lipoproteins

Elevated blood levels of cholesterol were recognized as a major risk factor for atherosclerotic diseases in the original Framingham studies [117]. Although LDL-C has been the principal target of preventive efforts through the widespread use of statins [118], other lipoprotein fractions, such as HDLs, are also important. Obesity in general is associated with increased serum triglycerides and decreased HDL-C [103]; the presence of insulin resistance is associated with an increase in this ratio [119]. The blood level of HDL-C is an important protective factor for atherosclerotic cardiovascular disease; thus, higher HDL-C is associated with a reduced risk of cardiovascular death, myocardial infarction, and stroke [120]. However, measures of HDL function may be better indices of cardiovascular risk than the common laboratory measure of the amount of cholesterol present in the HDL fraction [121]. The protective value of HDL may result from their ability to transport cholesterol from peripheral sites back to the liver, the so-called “reverse cholesterol transport” or “cholesterol scavenging” [122]. This function is particularly important at sites of inflammation, where excess cholesterol and other lipid molecules accumulate in macrophages [122]. Many factors can affect HDL function, and other mechanisms may contribute to their atheroprotective effects [123,124]; improvement in cholesterol scavenging by HDL is an active area for drug development [125,126].

Alterations in lipoproteins have been implicated as risk factors for ADs in some recent studies, although conflicting results have been reported. Thus, in one study, higher levels of HDL-C were associated with as much as a 50% to 60% reduction in the risk of ADs [127], whereas an opposite relationship was found in another study [128]. These apparent discrepancies may be attributable to differences in the method used for AD diagnosis (i.e., neuropathologic vs clinical) differences in subject populations (nursing home residents vs community-dwelling individuals) or other factors. Further research is needed to clarify these apparent discrepancies.

4.4.7. Inflammation

Obesity is associated with increased circulating levels of a number of cytokines known to reflect and promote inflammatory processes, such as tumor necrosis factor-α and interleukin-6 (IL-6) [129,130]. Some of these cytokines, such as IL-6, are produced by adipocytes; as much as 30% of circulating IL-6 can be accounted for by fat mass [131,132]. The expression and secretion of these proinflammatory cytokines [133] reflect, in part, their common, albeit remote, evolutionary origin with macrophages [104], with which they are heavily commingled in abdominal adipose tissue as indicated earlier in the text [109,110]. Recent research has shown that adipocytes produce a wide and complex array of mediators influencing inflammatory processes, insulin resistance, and energy metabolism [104,110,133], many of which have been implicated in the increased risk of atherosclerotic vascular diseases
associated with the metabolic syndrome [134]. However, the relative roles of these mediators in cardiovascular outcomes are far from clear as yet and remain an active area of investigation. Nevertheless, there is general agreement that low-grade systemic inflammation is an important pathophysiologic component of the accelerated atherosclerotic processes associated with metabolic syndrome [112,130,135,136].

4.4.8. Platelet activation and altered hemostatic functions

The vascular endothelial injury resulting from the atherosclerotic process results in the activation of platelet and hemostatic functions in several ways [137]. Intact function of the hemostatic system has a number of beneficial effects, including prevention of excessive bleeding and extravasation of vascular contents into tissue spaces, and vascular repair [138]; however, the cascade of events after sustained or unchecked activation of this system can have deleterious consequences [96,137,139]. In addition to the critical role of the vascular endothelium in actively maintaining a smooth surface to support laminar blood flow as described earlier in the text, the major components of the hemostatic system include platelets, the coagulation pathways, and fibrinolytic mechanisms [140]. All of these components have been implicated in the pathophysiology of atherosclerosis and thrombotic events [137,141,142] and in metabolic syndrome [143,144]. The critical role of platelet aggregation in acute coronary syndromes and cerebrovascular events forms the basis for the use of antiplatelet agents, such as aspirin and clopidogrel, in the prevention of such events, particularly in secondary prevention [145].

Two recent population-based longitudinal studies have shown an association between biomarkers of prothrombotic and/or antifibrinolytic activity and cognitive and functional decline [146] or the risk of developing VaD by NINDS-AIREN criteria [147]. Although many of the same biomarkers were assessed in these two studies, differences in outcomes make direct comparison difficult. Nevertheless, these findings provide support for the notion that a prothrombotic and/or antifibrinolytic state increases the risk of cognitive decline with aging.

4.4.9. CAA: Relationship with intracerebral vascular compromise and dementias

CAA refers to the deposition of Congo red staining material in or around cerebral blood vessels and/or leptomeninges on pathologic examination [148–152]. There are hereditary forms of CAA associated with single-gene mutations resulting in the production or overproduction of amyloidogenic proteins [150]; these include the hereditary forms of AD due to mutations in the amyloid precursor protein gene, although mutations in other amyloidogenic proteins have been described [152].

The most common forms of CAA are sporadic; these are associated with normal aging and are more common in individuals with dementias than in those without [151,152]. In a recent population-based neuropathologic study, moderate-to-severe CAA was found in 17 of 183 (10%) autopsies of individuals without dementia and in 81 of 243 (34%) autopsies of those with dementia [59]. In contrast to hereditary forms, in which several amyloidogenic proteins may be involved, the sporadic forms of CAA are almost always associated with deposition of amyloid precursor protein products, primarily β-amyloid40 [151,152]. It has been reported that as many as 80% to 100% of patients with AD examined at autopsy show evidence of CAA [151,152].

CAA is linked to the occurrence of intracerebral hemorrhages, including both lobar hemorrhages and cerebral microbleeds observed at autopsy [84]. This linkage is likely related to the accumulation of β-amyloid40 in perivascular spaces, resulting in the obstruction of the clearance of β-amyloid peptides from the CNS [84]. APOE genotypes APOE4 and APOE2 are associated with an increased presence and severity of CAA lesions and cerebrovascular hemorrhages [152].

4.5. Potential mechanisms by which vascular and metabolic disease processes convergently result in cognitive impairment

The human brain is heavily dependent on the integrity of its vascular supply owing to its high rate of oxidative metabolism, resulting in disproportionately high rates of glucose use and oxygen consumption for its relatively low mass [153]. In addition, the CNS lacks intrinsic stores of energy substrates, and is thus exquisitely dependent on the systemic circulation for continuous delivery of energy substrates [154]. The extensive tissue damage and functional impairment resulting from acute compromise of perfusion during an acute stroke is perhaps the clearest example of this intimate interdependency between local or regional blood flow and neuronal function. However, more subtle and insidious effects can occur over protracted time periods owing to progressive structural changes in cerebral blood vessels [155] and impairment in the normally closely regulated mutual coupling of blood flow and oxygen consumption [156]. It has been pointed out that lesser degrees of vascular compromise integrated over an extended time period may cause impairments in neuronal integrity and function similar to those caused by more abrupt and clinically observable ischemic events [157].

In Figure 1, we depict the mutually interacting metabolic and vascular factors that may contribute to dementia risk at a systemic level, as well as cellular and molecular mechanisms that have been proposed to underlie the structural damage to brain parenchyma that ultimately leads to cognitive and functional impairment. These cellular and molecular mechanisms are likely to include excitotoxicity due to excessive release of glutamate (albeit in a more localized manner than during acute stroke), alterations in intracellular calcium concentrations due to failure of intracellular calcium buffering mechanisms [158], blood–brain barrier
compromise [159], and activation of microglia and release of proinflammatory mediators [160]. Angiogenic factors are also released [161], although effective revascularization may be limited [162]. Hypoxia due to insufficient perfusion of and oxygen delivery to neuronal cells triggers a series of intracellular and intercellular responses, including activation of hypoxia inducible factor-1, mitochondrial dysfunction, and oxidative and nitrosative stress [163–165]. Ultimately, neuronal degeneration occurs through necrosis, apoptosis, and/or autophagy. Damage to oligodendrocytes and myelin also develops as a consequence of vascular compromise and the resulting cellular events [149]. Recent research suggests that vascular compromise interacts with the amyloid pathway in such a way as to exacerbate Alzheimer-type pathologic changes [159,163]; it has been proposed that cerebral microvessels are the initial nidus for the formation of amyloid lesions [166]. The reader is referred to several recent reviews that describe these events in greater detail [148,156,159,162,163,167,168]. However, many questions remain regarding the precise sequence of and interactions among these events, their relative importance in contributing to cognitive and functional impairment, whether and how these events can be detected through the use of biomarkers, and the most promising targets for interventions to prevent cognitive decline.

5. Recommendations for future research: Toward an integrated approach to the diagnosis, treatment, and prevention of dementias

We believe that several clear messages emerge from the above review of current research on the role of vascular diseases in the risk, course, and conceptualization of disorders characterized by aging-related cognitive impairment and decline.

5.1. Most dementias are associated with combinations of risk factors

The difficulties in categorizing dementias as AD, VaD, mixed dementia, and so forth are based on the assumption that these are distinguishable diseases. However, the available research shows that most cases of dementia are characterized by a mixed phenotype, both during life and at postmortem examination. The most common co-occurring conditions are those associated with AD and vascular dementias. In addition, many of the same risk factors predispose not only to dementias associated with plaques and tangles on postmortem brain examination but also to those associated with evidence of vascular disease. Thus, it would seem more pertinent to focus research efforts on elucidating the mechanisms by which vascular and neurodegenerative mechanisms jointly contribute to the development and progression of aging-related cognitive disorders than to consider them as separate diseases. In some respects, this integrative conceptual paradigm is analogous to the situation with the metabolic syndrome, in which multiple pathophysiologic mechanisms mutually interact to yield the clinical and pathologic outcomes of acute coronary syndromes, cerebrovascular events, and mortality.

5.2. Types of studies most likely to improve public health and advance knowledge

Based on our analysis of the current state of knowledge regarding vascular disease mechanisms in the risk,
dendantly evident from a recent analysis[171] estimating that to address these factors in the case of dementias is abun-
arable factors[170] that require distinct strategies. The need
practice [169]. The basis for this gap comprises multiple sep-
treatment do not always result in rapid changes in clinical
5.2.1. Translational and implementation research
Research findings regarding the effectiveness of a given
treatment do not always result in rapid changes in clinical
practice [169]. The basis for this gap comprises multiple sepa-
larable factors [170] that require distinct strategies. The need
to address these factors in the case of dementias is abund-
antly evident from a recent analysis[171] estimating that the
incidence of ADs could be dramatically reduced by a 10% reduction in the prevalence of major modifiable risk
factors, including several of those cited in the present article
(i.e., diabetes, hypertension, and obesity).

The emerging field of implementation science, also con-
sidered as a type of translational medicine[172], deals with the
barriers and obstacles that limit or delay the process of
modifying clinical care in response to the availability of sci-
centific evidence. Research, demonstration projects, and pro-
vider and consumer education are urgently needed to
translate current evidence about vascular disease prevention
and treatment into interventions that can mitigate risk or de-
lay the onset of disease. Importantly, such interventions
would have a broad impact on disease morbidity and mortal-
ity in the population owing to their high prevalence and the
multitude of adverse health consequences with which they are
associated.

5.2.2. Enhancing informational value of existing studies
There are a number of ongoing longitudinal studies of
clinical manifestations of dementias and cognitive decline, we propose here a blueprint for
achieving maximum benefits to health and society in ad-
dressing the urgent need to mitigate the impact of vascular
disease on cognitive function in susceptible populations.
We have organized this overall plan into (1) steps that can
(and should) be taken immediately based on our existing ev-
idence base, (2) strategies for enhancing the informational
value of currently active longitudinal research projects for
identifying key predictors of cognitive decline related to vas-
cular disease mechanisms, (3) recommendations for new re-
search initiatives to extend our knowledge base regarding
interactions among metabolic pathways and their ability to
predict future cognitive impairment and responses to treat-
ment, and (4) more long-range research strategies to make
maximum use of available data and suggest molecular and
systemic targets for the development of novel therapeutic
agents.

5.2.3. Designing new studies of midlife individuals to
examine the effects of earlier interventions on biomarkers of
adverse cognitive outcomes
There is typically a considerable lag time between the in-
ception of a longitudinal study, such as those currently in
progress, and the initiation of such studies, and an even lon-
ger latency for the emergence of data on key long-term out-
comes. Much new information is often accrued during this
time interval, which often cannot be readily obtained or
added once the study is in progress. As a result, there is a
need to design new studies incorporating current evidence
to take advantage of such advances. As an example, informa-
tion on multiple pathophysiologic domains associated with
insulin resistance, and perhaps some of the controversies and uncertainties that currently exist regarding their inter-
relationship, may be feasibly addressed by the initiation of
these studies assessing biomarkers representing multiple
domains contemporaneously over time. Some bio-
markers (e.g., those relating to platelet and hemostatic
system function) require specific sample collection or other
preanalytic methods that may not have been used in previous
studies of other kinds of biomarkers. Prospectively designed
studies would allow consideration of such factors before
study initiation, maximizing the quality of the data obtained.

5.2.4. Applying emerging principles, methods, and findings
of systems biology and network medicine to make optimal
use of existing knowledge and generate models of disease
processes that will lead to testable hypotheses about the
impact of interventions for which sufficient evidence does
not yet exist
Given the degree of interrelatedness among the systems
implicated in and influencing metabolic and vascular
dysfunction[173], a mathematical biologic approach can
substantially inform the elucidation of the relevant patho-
physiologic processes that contribute to the progression of the
disease process over time. For example, hypertension and T2DM are both identified earlier in the text as risk fac-
tors for vascular disease–mediated cognitive decline; how-
ever, hypertension is associated with insulin resistance,
which in turn is a risk factor for T2DM as well as for other
forms of vascular disease that may independently predispose
to cognitive impairment. Similarly, obesity is a risk factor for
both insulin resistance (and hence T2DM) and hypertension.
A mathematical biologic approach to quantifying the net im-
 pact of risk factors and their component pathophysiologic
mechanisms over time would integrate known data using sta-
tistical models of the dynamic and interactive processes

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A mathematical biologic approach to quantifying the net im-
 pact of risk factors and their component pathophysiologic
mechanisms over time would integrate known data using sta-
tistical models of the dynamic and interactive processes
involved, as well as identifying areas where data are incomplete and further research is needed.

Conceptualizing the disease process as a result of a loss of the ability to compensate for the cumulative burden of internal and external challenges faced over a lifetime [174] provides an opportunity to formulate key questions about the normal interregulatory relationships among these systems, that is, what relationships keep them from breaking down. Such a conceptual framework can help identify what parameters need to be measured not only to better define the critical nodal points influencing the expression of clinically and functionally significant cognitive impairment, but also to form testable hypotheses about where to target preventive and therapeutic interventions to achieve the maximum benefit to the individual.

6. Conclusions

Our understanding of the role of vascular disease processes in the development of aging-associated cognitive disorders has undergone several shifts over the past half-century. To some extent, the perceived importance of vascular disease to dementia risk was overshadowed for some time by the explosion of research on the underlying basis of the brain lesions identified by Alzheimer. This research has led to the identification of potential molecular targets for treatment and prevention, some of which are currently being tested. Nevertheless, it is critical to recognize that Alzheimer-type neuropathology usually does not occur in isolation, and vascular disease mechanisms are likely to play crucial roles in the evolution of the brain injury that eventually results in cognitive impairment and functional decline. We believe that a new paradigm is called for to advance research and mitigate the societal burden of these devastating diseases, one that is inclusive of both vascular and neuropathologic mechanisms. We outline a series of steps for applying this paradigm to current clinical practices and ongoing and future research.

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