Assessing the Preparedness of the Health Care System Infrastructure in Six European Countries for an Alzheimer's Treatment

KEY FINDINGS

- The burden of Alzheimer’s disease in high-income countries is expected to approximately double between 2015 and 2050. Recent clinical trial results give hope that a disease-modifying therapy might become available in the near future. The therapy is expected to treat early-stage patients to prevent or delay the progression to dementia.

- This preventive treatment paradigm implies the need to screen, diagnose, and treat a large population of patients with mild cognitive impairment. There would be many undiagnosed prevalent cases that would need to be addressed initially, and then the longer-term capacity to address incident cases would not need to be as high as the short-term capacity.

- We use a simulation model to assess the preparedness of the health care system infrastructure in six European countries—France, Germany, Italy, Spain, Sweden, and the United Kingdom—to evaluate, diagnose, and treat the expected number of patients.

- Projected peak wait times range from five months for treatment in Germany to 19 months for evaluation in France. The first years without wait times would be 2030 in Germany and 2033 in France, and 2042 in the United Kingdom and 2044 in Spain. Specialist capacity is the rate-limiting factor in France, the United Kingdom, and Spain, and treatment delivery capacity is an issue in most of the countries.

- If a disease-modifying therapy becomes available in 2020, we estimate the projected capacity constraints could result in over 1 million patients with mild cognitive impairment progressing to Alzheimer’s dementia while on wait-lists between 2020 and 2044 in these six countries.

- A combination of reimbursement, regulatory, and workforce planning policies, as well as innovation in diagnosis and treatment delivery, is needed to expand capacity and to ensure that available capacity is leveraged optimally to treat patients with early-stage Alzheimer’s disease.
Introduction

Alzheimer’s disease is a progressive neurodegenerative disorder that leads to cognitive and functional decline, resulting in Alzheimer’s dementia and premature death. Alzheimer’s dementia imposes a substantial burden on patients, their families and caregivers, and the broader society. As the risk of developing Alzheimer’s dementia increases with aging populations, the burden of disease is estimated to approximately double in G7 countries and nearly triple in G20 countries between 2015 and 2050 (Prince et al., 2015).

Currently, no disease-modifying treatment is available, but there are therapies in development. With some encouraging early-stage clinical trial results and collaboration efforts after many disappointing trials, there is cautious optimism that a therapy may become available in the near future (Aisen, 2017; "Alzheimer’s Disease: A Time for Cautious Optimism," 2015). Although there have been recent setbacks in trials that raise questions about the disease targets and mechanisms of action (Honig et al., 2018; Murphy, 2018), several therapies remain in clinical trials (Table 1).

Based on results from early-stage clinical trials, therapeutic development has focused on the hypothesis that Alzheimer’s dementia must be prevented rather than cured, because candidate treatments have not been able to reverse the course of dementia. Thus, current trials target patients with early-stage Alzheimer’s disease by first screening patients for signs of early-stage memory loss, or mild cognitive impairment (MCI), conducting tests for the Alzheimer’s disease pathology, and then treating patients with the aim of halting or slowing progression from MCI due to Alzheimer’s disease to Alzheimer’s dementia.

An important health systems challenge will arise if this new paradigm bears out in late-stage clinical trials. In the 28 European Union (EU) countries, we estimate that approximately 20 million individuals over age 55 have MCI (Eurostat, 2018; Petersen et al., 2018), although most people have not been tested for disease pathology, as there is currently no treatment option. Thus, when a therapy first becomes available, there would be a substantial number of existing (or prevalent) MCI cases that would require screening, diagnosis, and then treatment as quickly as possible to prevent the progression to full-blown Alzheimer’s dementia.1

We have previously analyzed the preparedness of the health care system in the United States to handle the potential caseload if and when a disease-modifying therapy becomes available (Liu et al., 2017). With the capacity of the current infrastructure projected forward, we found that there would be long waiting times for dementia specialist visits initially, followed by considerable waits for biomarker testing to confirm the Alzheimer’s pathology and moderate waits for infusion delivery of the therapy, resulting in 2.1 million patients developing Alzheimer’s dementia between 2020 and 2034 (about 13 percent of the potentially avoidable new cases in this period) while on waiting lists in the United States (Liu et al., 2017).

Other countries may face similar infrastructure challenges in diagnosing and delivering treatment to people with Alzheimer’s disease. International attention has been paid to the preparedness of individual countries to support and adopt innovation in dementia diagnosis, treatment, and care. For example, the Global Coalition on Aging and Alzheimer’s Disease International for G7 countries developed the Dementia Innovation Readiness Index (Global Coalition on Aging and Alzheimer’s Disease International, 2017). The World Health Organization has a global action plan with action area 4 focused on diagnosis, treatment, care and support (WHO, 2017), and a guide for countries to develop dementia plans (WHO, 2018).

The objective of this study is to extend the infrastructure analysis to six European countries: France, Germany, Italy, Spain, Sweden, and the United Kingdom. This selection is made up of six large countries representing 65 percent of the population in the European Union (Eurostat, 2018). As with the U.S. analysis, we assess how each country’s infrastructure for diagnosis and treatment lines up with the potential caseload of patients. We use the simulation model developed in the U.S. study and incorporate country-specific data and expert input on clinical practices. Our goal is to demonstrate the
potential magnitude of the mismatch between the patient caseload and the available supply of services, not to provide an exact prediction of future therapies and capacity. Our estimates reflect a base case scenario assuming that the current capacities of the health care systems are carried forward.

In the following sections, we briefly describe the conceptual framework and the simulation model. We then present data on the current capacity in each country. We show the effects of capacity constraints on access to diagnosis and treatment of Alzheimer’s disease and describe the effects of resolving the capacity constraints in each country. We conclude with a comparison of the preparedness of the health care systems in the six countries and discuss potential efforts to address the expected mismatch between supply and demand. We hope this analysis will help facilitate continued dialogue among stakeholders on how to ensure timely diagnosis and treatment of patients if a disease-modifying therapy for Alzheimer’s becomes available.

### Conceptual Framework and Simulation Model

The patient journey that we use as the framework for our analysis is depicted in Figure 1. We assume that individuals age 55 and older could undergo cognitive screening via a short instrument, such as the Mini–Mental State Examination or Folstein test in a primary care setting (screening phase) (Folstein, Folstein, and McHugh, 1975); we use the threshold
of 55 years based on the inclusion criteria in several late-stage clinical trials currently in progress (e.g., U.S. National Library of Medicine, 2018a; 2018b; 2018c). Patients screening positive would be referred to a dementia specialist for further evaluation and possible referral to testing for the presence of biomarkers related to Alzheimer’s disease (diagnostic phase). If testing is positive, patients would return to the dementia specialist for further evaluation to confirm whether treatment is indicated and appropriate. If appropriate, they would be referred to treatment (treatment phase), which would reduce the risk of progression from MCI to manifest dementia due to Alzheimer’s disease (outcomes phase). While patients await diagnosis and treatment, the disease continues to progress—that is, patients are at risk of progressing to a later stage of the disease, at which point the treatment would no longer be effective.

We use the simulation model previously developed for use in the United States to estimate the effect of capacity constraints on access to care for patients with suspected Alzheimer’s disease (Liu et al., 2017). The model consists of a Markov model to simulate transitions between disease states and a systems dynamic model to simulate health care system capacity constraints within the MCI state. Patients’ journey through the disease states is guided by transition probabilities from having “no cognitive impairment” (i.e., no MCI and no Alzheimer’s dementia) to MCI to Alzheimer’s dementia. The transition probabilities are based on
reported rates from epidemiological studies (see the appendix Table A.1). Within the MCI state, patients move through the diagnostic and treatment phases based on a system dynamics model with outflows constrained by infrastructure capacity. We model three capacity constraints: availability of dementia specialists, biomarker testing, and infusion delivery. We optimize the capacity of the dementia specialist workforce to provide the second (confirmatory) visit to a patient such that the maximum number of patients receive their confirmatory visit in the same year as their initial visit—that is, we assume that specialists would not take on a new patient for an initial visit if the specialist would not have the capacity to provide confirmatory visits for existing patients. Figure 2 shows the expected patient demand in the six countries at each stage of the patient journey, assuming no constraints on infrastructure capacity for screening, diagnosis, and treatment. We estimate that 7.1 million MCI patients would seek evaluation by a dementia specialist and 2.3 million would be indicated for treatment in the six countries. See appendix Figure A.1 for the expected patient demand in each country.

The assumptions about patient uptake, contraindications, and the treatment delivery and effectiveness are based on expert input in the U.S. study as described in Liu et al., 2017. For this analysis of the six European countries, we apply the same general assumptions regarding the treatment, uptake, and disease transitions as in our U.S. study and indicate when different assumptions are used. In summary, the key assumptions in our analysis are as follows:

- A disease-modifying therapy for patients with MCI due to Alzheimer’s disease becomes available in 2020. We assume the therapy would be an anti–beta-amyloid monoclonal antibody, as candidates that target beta-amyloid are the furthest along in clinical trials, that would be delivered by intravenous administration.

- Individuals become eligible for annual cognitive screening at age 55. Annual screenings are conducted by general practitioners, and we assume screening capacity would be unconstrained. We assume that 80 percent of eligible individuals would be screened each year. Screening starts in 2019 as patients and providers anticipate the approval of the therapy. We assume that 50 percent of individuals who screen positive for MCI would continue to receive further evaluation.

- Further evaluation would be conducted during a single visit to a dementia specialist—a neurologist, geriatrician, or geriatric (or old-age) psychiatrist, depending on the clinicians involved in dementia diagnosis and care in each country. If the evaluation confirms MCI and does not find an alternative explanation for MCI (e.g., prior stroke) or a reason to not pursue treatment (e.g., presence of another life-limiting disease), individuals would be referred to testing for biomarkers. We assume that 90 percent of patients with confirmed MCI that is possibly due to Alzheimer’s disease would undergo biomarker testing.

- In the European countries, biomarker testing may be performed by a cerebrospinal fluid (CSF) test or a Positron Emission Tomography (PET) scan for amyloid and/or
We assume that 90 percent of biomarker tests would be performed using CSF, and 10 percent would be conducted with PET scans when CSF is contraindicated (Alzheimer’s Research UK, 2018). We assume that 45 percent of MCI patients have clinically relevant biomarker levels.

- If a patient tests positive for biomarkers, she or he returns to a dementia specialist for a second visit, during which the treatment indication is confirmed, the patient is consented, and, if no contraindications to the treatment are found, the patient is referred to treatment. We assume that 80 percent of MCI patients testing positive for biomarkers would have no contraindications for treatment.

- We assume the following treatment characteristics: Treatment would be delivered by intravenous infusion every four weeks over one year, following protocols for a typical immunotherapy; and treatment would reduce the relative risk of progression from MCI to Alzheimer’s dementia by 50 percent.

For country-specific assumptions and data, we consulted a convenience sample of ten experts familiar with clinical practice and patient needs in each country—primarily clinicians and academic researchers specializing in care for patients with Alzheimer’s disease and related dementias. The country-specific assumptions include the type of specialists involved in detection and diagnosis of cognitive impairment and dementia and the use of CSF assays and PET imaging in the diagnosis of Alzheimer’s disease. The country-specific data include the population, disease prevalence, and mortality; the capacity scenarios are based on historical and projected workforce and infrastructure data. Table A.1 in the appendix contains the values for our model parameters and their respective sources.

Current Capacity Estimates and Projections

Specialist Workforce

We estimate the capacity of dementia specialists based on historical workforce data and assumptions about excess capacity. Overall, the dementia specialist workforce consists of neurologists, geriatricians, and geriatric or old-age psychiatrists. Table 2 shows the estimated number of specialists in each country.

As shown in Table 2, the specialists involved in diagnosing cognitive impairment and dementia vary in each country. We assume the types of specialties involved in this clinical area within each country to be unchanged over time in our projections. For example, diagnoses may be conducted by neurologists or geriatricians in memory clinics or private neurology practices in France. In Sweden, however, neurologists are typically not involved in diagnosis; rather, diagnoses are conducted by geriatricians and psychiatrists.

Another difference across countries is the extent to which psychiatrists have a role in formal diagnosis. In Germany and Sweden, experts suggested that approximately 60 percent of psychiatrists would be involved in formal diagnosis of MCI due to Alzheimer’s disease, whereas the remaining 40 percent primarily practice psychotherapy. In contrast, in the United Kingdom, old-age psychiatry is a relatively small subspecialty representing about 15 percent of total psychiatrists (Royal College of Psychiatrists, 2017).

Generally, there is a perceived shortage of specialists, which is expected to worsen with the aging populations, but country-specific workforce projections by specialty are scarce in publicly available reports. Publicly available workforce projections are limited to certain geographies, exhibit different trends by specialties, and show uncertainty in the projections. For example, the Centre for Workforce Intelligence in England estimated the total psychiatrist workforce in four possible scenarios, with a baseline projection of about a 1- to 2-percent decrease between 2015 and 2030 and alternate scenarios ranging from about a 9-percent decrease to a 5-percent increase in the psychiatrist
workforce (Centre for Workforce Intelligence, 2014). In Germany, the National Association of Statutory Health Insurance Physicians predicted the number of neurologists to remain relatively stable through 2030, with fluctuation of ±3 percent between 2014 and 2030 (Kassenärztliche Bundesvereinigung, 2016). In France, overall physician density is estimated to decrease slightly between 2015 and 2022 and then increase by over 25 percent by 2050 (Ono, Lafortune, and Schoenstein, 2013); these projections also include alternate scenarios but do not report on specific specialties. Given the variation in these workforce projections and the limited data, we use a simplified baseline assumption of static specialist workforce levels in each country from 2017 through 2050. If the workforce capacity were to change (e.g., if it responded to increased patient demand and expanded capacity), then our results would be different. Our base case scenario shows the effects if capacity is unchanged in the future. Therefore, our base case scenario should be interpreted as projections if the current workforce and capacity were maintained.

The availability of the dementia specialist workforce to see MCI patients in the diagnostic phase depends on the capacity of each specialist to conduct the evaluations. In our scenarios, we assume that each dementia specialist provides the same average number of ambulatory visits per year as neurologists in our U.S. model, 2,860 visits (Dall et al., 2013), because we assume the cognitive assessment and confirmation of the diagnosis would be conducted following the same guidelines across the specialist types. We also assume that these specialists devote 5 percent of their capacity to conducting these evaluations of MCI patients; we assume this excess capacity would be relatively low because of increasing demands on physician workforces with the aging population (we also examine scenarios using alternative assumptions with higher capacity, 7.5 percent, and lower capacity, 2.5 percent).

### Diagnostic Technology

Diagnosing predementia Alzheimer’s in a patient with MCI involves confirming the presence of biomarkers (beta-amyloid and/or tau) (Portet et al., 2006). In Europe, biomarkers may be detected using a CSF test or with neuroimaging. CSF samples are taken via a lumbar puncture (spinal tap) and tested for the presence of amyloid and tau proteins (Olsson et al., 2016; Herukka et al., 2017). Although lumbar punctures are more invasive than imaging tests, CSF tests are commonly used and generally well-accepted in the countries studied. However, if a patient has a contraindication for a lumbar puncture (e.g., due to anticoagulant medication) (Engelborghs et al., 2017), then neuroimaging with a PET scan may be used. In an amyloid PET scan, a radioactive tracer that selectively binds to amyloid is injected into a patient and a positron camera determines the brain areas with abnormal radiation activity, which indicates the presence of amyloid deposits.
We assume that capacity for CSF tests is theoretically unconstrained because lumbar punctures may be performed by general practitioners in various settings. We assume that 90 percent of biomarker tests could be conducted by CSF testing, and 10 percent would be conducted by PET scans (Alzheimer’s Research UK, 2018).

We project the capacity for conducting PET scans based on the historical number of PET scanners in each country and assumptions about the annual number of scans per scanner and growth in the number of PET scanners. Our projections are based on 2015 Organisation for Economic Co-operation and Development (OECD) data for an installed base of PET scanners in France, Italy, Sweden, and Spain (OECD, 2017b) and country-specific data sources for PET scanners in the United Kingdom and Germany (National Cancer Research Institute PET Core Lab, 2018; Gesundheitsberichterstattung des Bundes, 2018). As with our prior U.S. analysis, we assume that existing devices operate at about 50 percent of their capacity due to scheduling constraints. We assume that newly installed devices would be largely dedicated to amyloid scans and could be run at 80 percent of their capacity.

### Infusion Delivery

As mentioned, we assume the disease-modifying therapy would be administered intravenously. Thus, the capacity of the health care system to deliver infusions is a potential constraint to accessing the treatment. Many candidates currently in phase 3 trials are infused every four weeks over one to two years. However, this may reflect initial phases of treatment, and it is unknown whether longer-term durations of treatment would be needed. In this analysis, we assume that a course of treatment would consist of 14 infusions over one year. If the therapy required delivery for more than one year, then the capacity required would be even greater.

Our projections of the capacity of infusion centers are based on relative health care system capacity estimates and assumptions about growth rates. Due to lack of data on infusion capacity in the countries studied, we adjust the historical numbers of infusions delivered in the United States to each country based on population size and relative health care system infrastructure information. We develop a general health care capacity index for each country studied based on data on hospital beds, active nurses, MRI scanners, and PET scanners. The rationale for these four components is that the number of hospital beds is an indication of the facility space available, the number of nurses provides information about staff available to administer infusions, and the number of MRI and PET scanners provides an indication about medical technology and the degree of investment in the medical sector. We use OECD data to develop this index for the six countries relative to the United States (OECD, 2017b). We then use this index to scale the projected infusion capacity in each of the countries relative to the United States (based on infusion projections presented in Liu et al., 2017, and shown in appendix Figure A.2). Our infusion capacity index is shown in appendix Table A.2. We applied the same assumptions as in our prior U.S. analysis regarding the excess capacity (10 percent of current infusion capacity and 80 percent of new infusion capacity) that could be devoted to the treatment of MCI patients with Alzheimer’s pathology.

### Simulation Results for Selected Capacity Scenarios

We estimate the potential impact of capacity constraints on wait times and the progression of Alzheimer’s disease under three scenarios: a base case scenario with current capacity estimates projected forward, an alternative scenario without infusion capacity limits, and an alternative scenario without any capacity limits.

#### Base Case: Current Capacity Projections

Of the three constraints we consider, we estimate that there will be delays in dementia specialist visits and infusion treatment but find that biomarker testing is not a constraint in any of the countries studied. As shown in Figure 3, there are substantial differences in the capacity of the health care systems in the countries studied to handle a new treatment for
FIGURE 3
Projected Wait Times for Alzheimer’s Disease Diagnosis and Treatment, by Country
(average time delay in months)
The projected capacity for administering intravenous infusions in our base scenario falls short of potential demand.

Alzheimer’s disease. As a result, average wait times differ by country and are eliminated at different points in time: The backlog of cases clears as late as 2044 in Spain and as early as 2030 in Germany.

In France, wait times would be driven primarily by the availability of specialists. While wait times in France are eliminated within a few years, France has the highest capacity constraints in the short run due to the limited number of specialists (and, thus, longer average wait times that result in a higher share of avoidable Alzheimer’s dementia cases occurring compared with the other countries). The overall backlog clears by 2033, given the relatively larger infusion capacity in France.

In Spain and the United Kingdom, the wait times are a result of limited capacity for specialist visits and infusions. Because of the limited capacity in both the diagnosis and treatment phases, we estimate that the waiting period would extend to 2044 in Spain and 2042 in the United Kingdom.

In Italy, Sweden, and Germany, the wait times would be driven primarily by the capacity to deliver infusions. Wait times in Italy are expected until 2040. Wait times are much shorter in Sweden and Germany, where there are theoretically enough specialists to diagnose the large number of MCI patients. This results in wait times constrained only by availability of infusion services, which we expect to be relatively well-supplied compared with the other countries. We estimate that wait times extend until 2036 in Sweden and 2030 in Germany.

See appendix Figure A.3 for the number of patients waiting for diagnosis and treatment in each country.

The estimates for the base case scenario reflect what we believe are moderate assumptions about capacity growth; however, there is substantial uncertainty in these estimates given limited historical data and unknown characteristics of a future breakthrough Alzheimer’s therapy. In appendix Figure A.4, we show projections using alternate capacity assumptions that reflect different trajectories of growth to demonstrate a range of potential results given both higher and lower capacity assumptions for specialists and infusions. In the following section, we show projections of alternative scenarios assuming that policies are put into effect in response to a new therapy, specifically to address capacity issues. The alternative scenarios are discussed relative to the base case scenario.

Comparison with Alternative Scenarios

In addition to the base case scenario with the current capacity projected forward, we explore two alternative scenarios assuming the capacity constraints are partially or fully addressed. The healthcare system capacity for diagnosis and treatment could be expanded in response to a new Alzheimer’s treatment. We first discuss the specifications of the two alternative scenarios and then compare results of these scenarios for each country.

Alternative Scenario 1: Removing the Constraint on Infusion Capacity

For the first alternative scenario, we remove capacity constraints on infusion delivery in the simulation, leaving only access to dementia specialists
as a constraint, as biomarker testing is already not constrained in the base case scenario.

The projected capacity for administering intravenous infusions in our base scenario falls short of potential demand. As described earlier, we assume that patients would require infusions every four weeks for a period of one year. For an estimated 2.3 million MCI patients who could be eligible for treatment in 2020 in the six countries, approximately 32 million infusions would be required to treat the prevalent MCI population.

The historical growth in capacity for immunotherapy and chemotherapy treatments provides precedence for similar capacity growth to accommodate the demand. Similar challenges were addressed for immunomodulating antibodies for inflammatory diseases in the late 1990s, and, more recently, the introduction of biologic therapies for multiple cancers and other conditions, such as multiple sclerosis, have required expansion of infusion capacity (Tralongo et al., 2011).

It is possible that dementia specialists and other operators could also expand capacity in response to demand in a similar way; however, accommodating the large number of patients with MCI due to Alzheimer’s disease could be more challenging than treating patients with inflammatory conditions. An Alzheimer’s disease–modifying therapy would be first in class, whereas the immunomodulating antibodies were preceded by disease-modifying oral drugs, which allowed physicians to first target the limited infusion capacity to refractory cases. In addition, building infusion capacity for the immunotherapies was more financially viable because they are delivered to patients for the rest of their lives, while the amyloid antibodies are expected to be administered in a finite number of doses. Permanent capacity growth for amyloid antibodies may be less favorable, given that there will be about 14 million prevalent MCI cases initially but about 1 million incident MCI cases each year in the six countries studied.

An option to expand capacity quickly, in addition to infusion centers as extensions of memory clinics or other facilities, would be home infusion delivery. Home infusion services would reduce investment into permanent facilities that may later have idle capacity if the therapy is time-limited. The amyloid antibodies are infused over a relatively short time and are not known to have acute side effects; thus, they may be suitable for home infusion if it proves to be a financially viable option.

**Alternative Scenario 2: Removing All Capacity Constraints**

In alternative scenario 2, we remove all capacity constraints—that is, the constraints on both infusion delivery and dementia specialist visits. As with the prior scenarios, this is hypothetical; however, this scenario is less likely because medical specialists require long training times, and diagnosing Alzheimer’s requires comprehensive assessment. The assessment, including medical history, cognitive testing, neurological testing, and brain imaging, is typically conducted by physicians. We include this scenario to demonstrate the upper bound of Alzheimer’s dementia cases that could be potentially avoided if all capacity constraints are overcome.

**Comparison of New Alzheimer’s Dementia Cases Occurring in the Base Case and Alternative Scenarios**

Figure 4 shows how the base case scenario and the two alternative scenarios would change the estimated cumulative incidence of Alzheimer’s dementia between 2020 and 2050. Although the incremental benefit of removing capacity constraints is a small proportion of total possible incident cases avoided, it represents over 1 million additional patients in the six countries studied—a significant burden for the respective health care systems.

Between 2020 and 2050, we estimate that over 3.9 million incident Alzheimer’s dementia cases would be avoided in the base case in France.7 Removing the capacity constraint for infusion delivery (Scenario 1) would avoid an additional 32,000 cases relative to the base case (0.7 percent of potentially avoidable new cases). Removing the specialist constraint as well (Scenario 2) has a larger impact and would avoid 389,000 additional cases relative to the base case (9.0 percent of potentially avoidable new cases).

In the United Kingdom, nearly 3.7 million incident Alzheimer’s dementia cases would be avoided in
NOTES: The number of cases in both Scenario 1 and Scenario 2 are shown relative to the base case scenario. For example, in Germany, there are 55,000 additional new Alzheimer’s dementia cases averted in Scenario 1 relative to the base scenario, and 55,000 additional cases averted in Scenario 2 relative to the base scenario. In other words, there is no difference between Scenario 1 and Scenario 2, in which the specialist constraint is removed, because there is no projected waiting time for specialists in Germany.
the base case. We estimate that removing the capacity constraint for infusion delivery would avoid an additional 89,000 cases relative to the base case (2.2 percent of potentially avoidable new cases), and when the specialist constraint (Scenario 2) is removed, an additional 260,000 cases would be avoided relative to the base case (6.4 percent of potentially avoidable new cases).

In Spain, we estimate that nearly 3 million incident Alzheimer’s dementia cases could be avoided in the base case. Removing the capacity constraint for infusion delivery (Scenario 1) would allow an additional 83,000 cases to be avoided relative to the base case (2.6 percent of potentially avoidable new cases). When the specialist constraint (Scenario 2) is removed, an additional 171,000 cases would be avoided relative to the base case (5.3 percent of potentially avoidable new cases).

In Italy, we estimate over 4.3 million incident Alzheimer’s dementia cases would be avoided in the base case, an additional 101,000 cases relative to the base case (2.2 percent of potentially avoidable new cases) would be avoided if the capacity constraint for infusion delivery is removed, and 146,000 additional cases would be avoided relative to the base case (3.2 percent of potentially avoidable new cases) if the specialist and infusion constraints are removed.

In Sweden, over 600,000 incident Alzheimer’s dementia cases would be avoided in the base case. Removing the capacity constraint for infusion delivery (Scenario 1) would allow an additional 11,000 cases to be delayed relative to the base case (1.7 percent of potentially avoidable new cases). When the specialist constraint (Scenario 2) is removed, we estimate that 12,000 additional cases would be avoided relative to the base case (1.9 percent of potentially avoidable new cases) between 2020 and 2050.

In Germany, over 5.6 million incident Alzheimer’s dementia cases would be avoided in the base case scenario. Removing the capacity constraint for infusion delivery (Scenario 1) would allow an additional 55,000 cases to be avoided relative to the base case (1.0 percent of potentially avoidable new cases). There are no additional avoided cases when the specialist constraint (Scenario 2) is removed relative to the base case because we estimate no waiting times for specialist visits even in the base case, hence the number of avoidable cases relative to the base scenario is also 55,000.

Summary of Results by Country

Table 3 shows a summary of the differences in maximum wait times across the six countries. The longest wait times are experienced in the first few years after a new therapy becomes available. As noted

<table>
<thead>
<tr>
<th>Country</th>
<th>Maximum Waiting Time, Months</th>
<th>First Year with No Wait Times</th>
<th>Potentially Avoidable New Alzheimer’s Dementia Cases Occurring While Patients Wait, 2020–2044 (percentage of potentially avoidable cases)</th>
<th>Base Case: Specialist and Infusion Constraints</th>
<th>Scenario 1: Specialist Constraint</th>
</tr>
</thead>
<tbody>
<tr>
<td>Germany</td>
<td>No wait &lt; 6</td>
<td>2030</td>
<td>55,000 (1%)</td>
<td>0 (0%)</td>
<td></td>
</tr>
<tr>
<td>Sweden</td>
<td>&lt; 6 6–12</td>
<td>2036</td>
<td>12,000 (2%)</td>
<td>1,000 (&lt; 1%)</td>
<td></td>
</tr>
<tr>
<td>Italy</td>
<td>&lt; 6 6–12</td>
<td>2040</td>
<td>146,000 (3%)</td>
<td>45,000 (1%)</td>
<td></td>
</tr>
<tr>
<td>Spain</td>
<td>6–12 6–12</td>
<td>2044</td>
<td>171,000 (5%)</td>
<td>88,000 (3%)</td>
<td></td>
</tr>
<tr>
<td>United Kingdom</td>
<td>&gt; 12 6–12</td>
<td>2042</td>
<td>260,000 (7%)</td>
<td>171,000 (4%)</td>
<td></td>
</tr>
<tr>
<td>France</td>
<td>&gt; 12 &lt; 6</td>
<td>2033</td>
<td>389,000 (9%)</td>
<td>357,000 (8%)</td>
<td></td>
</tr>
<tr>
<td>Total</td>
<td>—</td>
<td>—</td>
<td>1,033,000 (5%)</td>
<td>662,000 (3%)</td>
<td></td>
</tr>
</tbody>
</table>

NOTE: The potentially avoidable new Alzheimer’s dementia cases are measured relative to Scenario 2, in which there are no capacity constraints and, thus, no patients waiting. There are no waiting times for biomarker testing if SCF assays are used for 90 percent of testing and PET scans are used for 10 percent of testing.
earlier, we find that biomarker testing is not a binding constraint, given the availability of CSF testing in Europe. All six countries are expected to experience infusion wait times less than 12 months, with infusion wait times of less than six months estimated in Germany and France. There is greater variation in specialist wait times, with no wait times expected in Germany and wait times more than 12 months expected in France and the United Kingdom.

Across the six countries, between 1 and 9 percent of potentially avoidable incident cases of Alzheimer’s dementia between 2020 and 2044 still occur in the base scenarios (using Scenario 2, without any constraints, as the reference point for the maximum number of avoidable cases). The percentage of avoidable cases is highly dependent on the effectiveness of the therapy; however, the relative differences between countries provide insight on the variation in the infrastructure across countries. The smallest constraint is observed in Germany, with 99 percent of cases avoided already in the base scenario. The elimination of wait times for specialists would have the largest relative impact for Spain, the United Kingdom, and France. Although the vast majority of new Alzheimer's dementia cases are avoided in the base scenarios in all countries, over 1 million cases develop while patients are on waiting lists in the six countries studied. If infusion capacity can be addressed (Scenario 1), the number of potentially avoidable new Alzheimer’s dementia cases that occur while patients are on wait lists goes down to 662,000. However, the specialist capacity must be addressed to prevent waits for all possible new dementia cases.

**Limitations**

There are several limitations in our analysis. As with our prior analysis, the framework for the model is based on a highly stylized patient journey that simplifies patterns of care. We model hypothetical scenarios involving a future Alzheimer's treatment with assumed specifications—such as time of market entry, efficacy, and dosing—that depends on actual clinical trial results and is uncertain at this time. Nonetheless, the goal of this analysis is to describe possible scenarios rather than precisely predict future supply and demand to demonstrate potential challenges and inform policy solutions that could help mitigate the challenges.

Many of our modeling assumptions are based on a therapeutic profile that is unknown at this time. As current clinical trials of Alzheimer’s therapies typically include patients age 50 or 55 and older (e.g., U.S. National Library of Medicine, 2018a; 2018b; 2018c), we assume that patients would be screened starting at age 55. We assume uptake rates for screening, evaluation, testing, and treatment based on expert input from our prior U.S. analyses; however, uptake at each step would depend on such factors as patient awareness and acceptance and reimbursement levels in each country.

Our capacity projections are based on extrapolations of historical data and assumptions informed by expert input from clinicians and researchers familiar with the countries studied, and our projections will almost certainly not accurately predict future patterns. Although general practitioners have the technical capability to do basic cognitive screening and lumbar punctures (to collect cerebrospinal fluid for diagnostic tests), not all general practitioners may choose to perform these tests. We do not examine the impact of constraints in primary care settings; any capacity constraints in cognitive screening and lumbar punctures would increase estimated waiting times. While CSF and PET measures are correlated and both can accurately detect early Alzheimer's disease, the tests provide different information about disease state and progression (Palmqvist et al, 2015; Bouallegue et al., 2017; AlzForum, 2017); we did not differentiate between CSF and PET testing in this analysis. We also do not consider capacity issues related to monitoring the disease progression, such as the use of magnetic resonance imaging.

Due to the lack of data on the specialist workforce in the future and on current capacity to provide infusions in each of the countries studied, we make simplified assumptions with a flat workforce projection and based on relative capacity data compared with the United States. We do not distinguish subspecialties within neurology; some general neurologists may require training in dementia diagnosis and care. We also assume that specialists across countries see the same average number of patients per year.
We estimate that the combined effect of the specialist and infusion wait times could result in more than 1 million new cases of Alzheimer’s dementia that could be avoided between 2020 and 2044.

Although the number of consultations per doctor varies across countries, the number of doctor consultations depends partly on the role of nurses and other practitioners in providing care (OECD, 2017a). We assume that early detection and diagnosis protocols for cognitive impairment would require a similar amount of time regardless of the country. However, our convenience sample of clinical and patient advocacy experts did not include specific expertise in workforce or infusion market issues. Because of these potential shortcomings, we include projections of alternative capacity assumptions in appendix Figure A.4 to show a range of potential results, given both higher and lower capacity assumptions for both specialists and infusions.

As with most simulation analyses, we combined data from various sources that may not be consistent with each other. For example, there are multiple sources for dementia and Alzheimer’s dementia prevalence estimates. We relied on country-specific estimates over international estimates when possible and verified that our model produces similar projected burden of Alzheimer’s dementia as other studies (see appendix Figure A.5). In addition, we make several simplifications, such as transitions between disease states occurring in one-year time steps and parameterized by annual transition probabilities. As developments in diagnostic and treatment capacity depend on a host of uncertain factors, such as the results of clinical trials, level of demand, and reimbursement levels, we emphasize that our results are meant to show the magnitude of the mismatch between projected demand and capacity rather than to provide a precise estimate of that mismatch.

Discussion

There is guarded optimism among researchers that one or more disease-modifying therapies for Alzheimer’s disease could become available in the upcoming years. Such a breakthrough therapy could finally provide patients and their families with a treatment to delay or prevent the progression of this devastating condition. The potential caseload is sizable, given that most therapies in development focus on treating early-stage Alzheimer’s disease: We estimate that approximately 7.1 million people with MCI in the six European countries could seek timely diagnosis by a specialist and, when indicated, testing for Alzheimer’s pathology to determine their eligibility for treatment.

Our analysis suggests that the health care systems in some of the European countries have insufficient capacity to diagnose and treat the large number of patients with early-stage Alzheimer’s disease. While there would be no waiting for biomarker testing if capacity for CSF testing is sufficient, we estimate that all countries could face insufficient capacity for infusion services, and some countries also have limited availability of specialists under our base case capacity assumptions. In Germany and Sweden, the main infrastructure constraint would be infusion capacity, resulting in less than 2 percent of potentially avoidable new Alzheimer’s dementia cases developing in each country while patients wait. However, in the other four countries, wait times due to both specialist availability and infusion capacity would delay treatment for more significant numbers of patients, resulting in up to about 9 percent
of potentially avoidable Alzheimer’s dementia cases while patients are waiting in France, 6 percent in the United Kingdom, 5 percent in Spain, and 3 percent in Italy. As Alzheimer’s disease is a progressive neurodegenerative disorder, we estimate that the combined effect of the specialist and infusion wait times could result in over 1 million new cases of Alzheimer’s dementia that could be avoided between 2020 and 2044, were a treatment to exist and be delivered immediately to all eligible patients.

We emphasize that our projections assess the potential capacity to deliver an Alzheimer’s therapy, whereas regulations, reimbursement levels, and volume agreements between regulators, payers, and providers will determine the actual capacity devoted. In addition, the projections hinge on assumptions that are uncertain, given limited information on how capacity will evolve over time. For example, resource allocation in the National Health Service (NHS) of the United Kingdom is determined by centrally mandated requirements and decisions by clinicians in the local NHS provider organizations. In Germany, capacity planning and volume agreements at the state and regional levels will influence available capacity and payment levels, which, in turn, will affect the degree to which providers will devote activity to dementia-related care.

Causes of Differing Wait Times Between Countries

The different patterns in wait times are driven by the relative capacity for specialist visits and infusion delivery in each health care system. Notably, the limited capacity for both specialists and infusion delivery in Spain and the United Kingdom resulted in the longest period before wait times are eliminated (i.e., until the backlog of cases is cleared). However, the peak wait times from diagnosis to treatment were the longest in France, due to the substantial wait times for specialist visits in the early years. In contrast, we estimate wait times for infusions but minor wait times for specialists in Italy. While capacity for infusion delivery results in delayed access to treatment in Sweden and Germany, overall wait times are much shorter there than in the other countries.

The range of specialties involved in dementia care is the critical factor in determining wait times for specialist visits. Psychiatry is a much larger specialty than neurology and geriatrics in all six countries. Thus, countries like Germany and Sweden, where a larger share of psychiatrists is involved, tend to have short wait times. Conversely, wait times are long in countries, like the United Kingdom and France, where fewer or no psychiatrists are involved.

In countries that currently rely on the relatively small neurology and geriatric specialties to conduct the diagnosis and evaluation of early-stage Alzheimer’s patients, it may be possible to expand the role of physicians from larger specialties. For example, more internal medicine physicians and general psychiatrists could be trained in Alzheimer’s diagnosis. Such training and coaching by specialists may be complemented by telemedicine and telemonitoring efforts currently under way in many European countries (eHealth Stakeholder Group on Implementing the Digital Agenda for Europe, 2014).

Some Countries Have Initiated Efforts to Improve Access to Dementia Care

In addition to training more providers, the establishment of graduated clinical pathways that rely

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**Box 1. Plan Maladies Neurodégénératives 2014–2019, France**

The French Ministers of Health, Ageing and Family and of Research have published a five-year strategic plan to improve care for neurodegenerative disorders (Alzheimer Europe, 2016). The plan emphasizes the need for care pathways for these disorders generally, and specifically for Alzheimer’s disease. It points out the need for screening and early detection, proper diagnostic evaluation, and management of comorbidities, even in the absence of a disease-modifying treatment. The plan calls for the development of better tools for screening and diagnosis, access to lifelong quality care, wherever patients live, and the need to build interdisciplinary capacity and capabilities, mainly through centers of excellence and diffusion into the community.
on general practitioners to screen and identify higher-risk patients out of the large pool of prevalent MCI patients could relieve the burden on specialists. For example, France has a national plan that includes aims to improve care pathways for patients with neurodegenerative disorders (see Box 1). Italy has launched a project to identify patients with the highest risk of developing Alzheimer’s dementia to improve coordination and timeliness of care (see Box 2). An understanding of the characteristics of patients with early-stage Alzheimer’s disease is vital to efficiently identify patients who could benefit from a new treatment.

Other Countries Have Explored New Models for Infusion Delivery

If a therapeutic agent requires near-monthly visits to administer an intravenous infusion, most European countries may not have the capacity to provide infusions to all eligible patients in a timely manner. In contrast to the specialist limitations, the inadequacy of infusion capacity may have two relatively near-term solutions: building dedicated infusion centers as extensions to existing facilities and using home infusions, if allowable based on the therapy’s safety profile.

Dedicated outpatient infusion centers may be built in cooperation with hospitals, memory clinics, and other health care facilities. In the past, specialized infusion centers have been established to provide outpatient infusion therapy administered by nurses to multiple sclerosis patients in Germany (see Box 3). Similar centers could be established for MCI patients if a new therapy becomes available, with specialized nurses trained in intravenous administration, oversight by physicians, and patient education.

Although there is currently limited experience with home infusions in Europe, a mix of facility-based and home infusions may ultimately offer patients and physicians therapeutic options when resources are constrained, assuming both are

Box 2. Interceptor Project, Italy

In December 2017, Italy launched the Interceptor Project to help identify people at higher risk of developing Alzheimer’s disease, with the goal of improving their quality of care (Alzheimer Europe, 2017). The initiative aims to enroll 400 patients ages 50 to 85 with mild cognitive impairment and develop strategies to identify patients with the highest risk of developing Alzheimer’s dementia and thus the greatest likelihood of benefiting if a new treatment becomes available (“Italy Launches Pioneering Project to Identify Alzheimer’s Risk,” 2017). The Italian Ministry of Health expects to collect comprehensive data based on neuropsychological assessments, CSF tests, and PET and MRI scans over a period of three years, which is expected to help inform the selection of the most appropriate biomarkers for nationwide screening for patients with a higher risk of progressing toward Alzheimer’s dementia. One of the stated goals, moreover, is to better understand which patients are most likely to benefit from a novel therapy, thus increasing the system’s sustainability (Ministero della Salute, 2017).

Box 3. Multiple Sclerosis Specialist Nurses in Germany and Other EU Countries

The advent of new therapies for multiple sclerosis, a severe neurological condition, has necessitated new drug delivery pathways, including the education of nurses to provide specialized care for patients with regular antibody infusions. In Germany, it is possible to receive infusions for multiple sclerosis therapy in outpatient settings with specialized nurse assistance, increasing the quality of life of patients while reducing the cost and burden on existing health care facilities in caring for these patients (Giedigkeit, 2018).

Where relevant, nurses may also provide training for self-administered injections and the education about potential side effects (Sächsisches Krankenhaus Großschweidnitz, 2018). Across European countries, an online training program for nurses that aims to consolidate standards in multiple sclerosis care has been offered in multiple languages and offers six different modules: understanding the disease, clinical presentation, diagnosis and assessment, treatment, care and support, and rehabilitation (European Multiple Sclerosis Platform, 2018).
implemented in compliance with relevant regulations and are found to be financially viable.

EU-wide Efforts Can Provide Guidance and Promulgate Best Practices

In addition to increasing specialist and infusion capacity in each country, joint actions and planning in Europe can help provide better coordinated and more timely care for Alzheimer’s patients. For example, the Second European Joint Action on Dementia aims to identify best practices in dementia care (see Box 4), and several national plans and strategies have been drafted in countries including France, Italy, and Sweden to improve the diagnosis and care of dementia patients, improve their quality of life (and that of their caregivers), and better coordinate research (Alzheimer Europe, 2016; Di Fiandra, 2015; Alzheimer Europe, 2013).

Box 4. European Joint Action on Dementia

The second European Joint Action on Dementia was launched in 2016, building on a previous joint action ALCOVE (Alzheimer Cooperative Valuation in Europe) that sought to improve epidemiological data on dementia, improve the timeliness and reliability of dementia diagnosis, understand support systems for behavioral and psychological symptoms, improve the rights and dignity of dementia patients, and reduce the use of antipsychotics to improve dementia patients’ quality of life (Alzheimer Europe, 2018a; Leperre Desplanques et al., 2013).

The most recent effort has brought together experts from multiple European countries with the goal of discussing “practical guidance for policy makers developing and implementing their national dementia plans, policies and strategies” (Alzheimer Europe, 2018a). In November 2017, for instance, they published a report on the risks and benefits associated with dementia diagnosis (Krolak-Salmon et al., 2017). In early 2018, a pilot program addressing diagnosis and post-diagnostic support was launched, focusing on destigmatization, nurse–general practitioner cooperation, and telemedicine to improve dementia detection in nursing homes (Act on Dementia, 2018).

Additionally, the Joint Action on Dementia aims to generate evidence on improving outcomes in dementia patients and their caregivers; enhance EU-wide collaboration on dementia diagnosis, services, and support; and inform an international collaboration on care for dementia patients (Alzheimer Europe, 2018b).

Conclusions

Positive early clinical trial results have led to guarded optimism that a disease-modifying therapy for Alzheimer’s disease may become available in the coming years. The ability to halt or slow the progression of this devastating and common disease would represent a significant breakthrough. We find variation in the preparedness of the health care systems in six European countries studied: All six systems may encounter wait times due to lack of capacity, but some systems have less infrastructure than others to diagnose and treat early-stage Alzheimer’s patients. Our analysis suggests that the health care systems in some European countries would lack adequate capacity to provide patients with access to treatment within a reasonable time frame, mainly because of constraints in the capacity of specialists to diagnose patients and of infusion delivery services. If a disease-modifying therapy becomes available in 2020, we estimate that failure to increase capacity means that over 1 million patients with MCI due to Alzheimer’s disease could develop dementia while waiting for evaluation and treatment in the six countries between 2020 and 2044. Longer wait times and thus higher shares of potentially avoidable dementia cases are expected for France, the United Kingdom, and Spain.

Addressing the health care capacity constraints will require combining payment, regulatory, and workforce planning policies with broad education campaigns to increase patient awareness of the importance of timely detection of early-stage disease. We hope that this report helps facilitate discussions among multiple stakeholders on how to address the potential obstacles to delivering an Alzheimer’s disease therapy in a timely way.


OECD—See Organisation for Economic Co-operation and Development.


WHO—See World Health Organization.


———, A Study of CAD106 and CNP520 Versus Placebo in Participants at Risk for the Onset of Clinical Symptoms of Alzheimer’s Disease (Generation S1), National Institutes of Health, U.S. Department of Health and Human Services, 2018c. As of May 25, 2018: https://clinicaltrials.gov/ct2/show/NCT02565511
The caseload would lessen after the prevalent cases are treated, as the number of patients who newly develop MCI (incident cases) each year would be a fraction of the prevalent cases. In this study, we focus on cases in six European countries, in which we estimate approximately 14.3 million individuals over age 55 currently have MCI and about 1.0 to 1.4 million incident cases of MCI will develop each year between 2020 and 2050, suggesting the long-term capacity would not need to be as high as the short-term capacity needed when a new treatment is introduced.

This assumption differs from our prior analysis in the United States, where a PET scan is the only currently Food and Drug Administration–approved modality for clinical use.

Although we assume that 90 percent of biomarker tests could be conducted using CSF in each of the six countries in this analysis, there is within-country regional variation in the acceptability of using CSF for diagnostic testing.

The evaluation of MCI patients may include a brain magnetic resonance imaging (MRI), which we do not assume to be constrained by capacity. Although the average number of consultations per doctor varies across countries, cognitive assessment and diagnosis following standardized protocols could reduce the variation in length of visits. If fewer consultations per doctor are possible, then the capacity projections would be lower; see appendix Figure A.4 for an alternate low-capacity scenario.

Although our convenience sample of experts from the six countries confirmed that general practitioners are trained to perform lumbar punctures in various settings, there is variability in acceptance and norms across and within countries.

The actual treatment duration would depend on the results from later-stage clinical trials.

In our scenarios, we assume treatment reduces the relative risk of progression from MCI to Alzheimer’s dementia by 50 percent. We refer to the difference in Alzheimer’s dementia cases that occur with the treatment without any infrastructure constraints and those that occur without the treatment as “potentially avoidable” cases within a given time period. Although we refer to these cases as avoidable, in reality it is unknown whether a treatment would delay the progression of Alzheimer’s disease for some time or prevent dementia from occurring.
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About This Report

This report shows the potential magnitude of health care infrastructure constraints in the evaluation and treatment of early-stage Alzheimer’s disease with a future disease-modifying therapy in six European countries. This research was sponsored by Biogen and conducted within RAND Health, a division of the RAND Corporation. A profile of RAND Health, abstracts of its publications, and ordering information can be found at www.rand.org/health. The authors thank the following subject-matter experts for sharing their insights into clinical practices and patient needs in the six countries: Antonio del Olmo, Lutz Frölich, Michael Karran, Pierre Krolak-Salmon, José Luis Molinuevo, Matthew Norton, Patrizia Spadin, Anders Wimo, and Bengt Winblad. The authors also thank Jon Sussex and Peter Hudomiet for their valuable feedback on this report.