Venetoclax Combined With Low-Dose Cytarabine for Previously Untreated Patients With Acute Myeloid Leukemia: Results From a Phase Ib/II Study

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abstract

PURPOSE Effective treatment options are limited for patients with acute myeloid leukemia (AML) who cannot tolerate intensive chemotherapy. An international phase Ib/II study evaluated the safety and preliminary efficacy of venetoclax, a selective B-cell leukemia/lymphoma-2 inhibitor, together with low-dose cytarabine (LDAC) in older adults with AML.

PATIENTS AND METHODS Adults 60 years or older with previously untreated AML ineligible for intensive chemotherapy were enrolled. Prior treatment of myelodysplastic syndrome, including hypomethylating agents (HMA), was permitted. Eighty-two patients were treated at the recommended phase II dose: venetoclax 600 mg per day orally in 28-day cycles, with LDAC (20 mg/m² per day) administered subcutaneously on days 1 to 10. Key end points were tolerability, safety, response rates, duration of response (DOR), and overall survival (OS).

RESULTS Median age was 74 years (range, 63 to 90 years), 49% had secondary AML, 29% had prior HMA treatment, and 32% had poor-risk cytogenetic features. Common grade 3 or greater adverse events were febrile neutropenia (42%), thrombocytopenia (38%), and WBC count decreased (34%). Early (30-day) mortality was 6%. Fifty-four percent achieved complete remission (CR)/CR with incomplete blood count recovery (median time to first response, 1.4 months). The median OS was 10.1 months (95% CI, 5.7 to 14.2), and median DOR was 8.1 months (95% CI, 5.3 to 14.9 months). Among patients without prior HMA exposure, CR/CR with incomplete blood count recovery was achieved in 62%, median DOR was 14.8 months (95% CI, 5.5 months to not reached), and median OS was 13.5 months (95% CI, 7.0 to 18.4 months).

CONCLUSION Venetoclax plus LDAC has a manageable safety profile, producing rapid and durable remissions in older adults with AML ineligible for intensive chemotherapy. High remission rate and low early mortality combined with rapid and durable remission make venetoclax and LDAC an attractive and novel treatment for older adults not suitable for intensive chemotherapy.

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INTRODUCTION The median age at diagnosis of acute myeloid leukemia (AML) is 68 years. Older adults are often ineligible for intensive chemotherapy and thus have limited effective treatment options. Less-intensive approaches to treatment, such as low-dose cytarabine (LDAC), are associated with poor response rates (11% to 19%) and median survival times (≤6 months). Similarly, initial treatment with azacitidine, decitabine, or gemtuzumab ozogamicin result in complete remission (CR) plus CR with incomplete blood count recovery (CRi) rates of less than 30%.

patients do not receive leukemia therapy. These factors underscore the high unmet need for more-effective and less-toxic treatment options for older adults with AML, particularly those who are ineligible for intensive chemotherapy.

B-cell leukemia/lymphoma-2 (BCL-2) family members, including BCL-2, BCL-XL, and MCL1, promote cell survival by binding and sequestering pro-apoptotic proteins in cancer cells. BCL-2 has been shown to mediate chemoresistance and enhance survival of leukemic blast and progenitor cells. Venetoclax, a selective, potent, orally bioavailable small-molecule BCL-2 inhibitor, has been studied alone and in combination with other agents.
combination with other agents in several hematologic malignancies. A phase II study reported an overall response rate (ORR) of 19% with venetoclax monotherapy in heavily pretreated patients with AML. Resistance to venetoclax monotherapy may be mediated by other pro-survival proteins, such as MCL1, that sequester endogenous BCL-2 homology domain 3-only proteins (eg, Bim) released by venetoclax on BCL-2 binding. Several drugs—including anthracyclines, hypomethylating agents (HMAs), and cytarabine—have demonstrated the ability to down-regulate MCL1 expression and act synergistically with venetoclax against AML cells in preclinical studies. As proof of concept, a 61% CR/CRI rate was reported for venetoclax combined with HMAs (ie, azacitidine or decitabine) in treatment-naïve older adults with AML, exceeding previously reported response rates for HMAs alone. Here, a phase Ib/II study was conducted to determine the safety and preliminary efficacy of venetoclax in combination with LDAC in previously untreated adults with AML who were ineligible for intensive chemotherapy.

**Patients and Methods**

**Patients**

Patients age 60 years or older with previously untreated AML and ineligible for intensive chemotherapy were enrolled (Data Supplement). Patients with secondary AML or prior treatment with HMAs for myelodysplastic syndrome (MDS) were permitted. Exclusion criteria included prior therapy for AML or any previous use of cytarabine for any indication (more details in the Treatment section). Local ethics committee approval was obtained, and patients provided written informed consent. The study was conducted in accordance with the International Conference on Harmonization, Good Clinical Practice guidelines, and the Declaration of Helsinki.

**Study Design**

This open-label, multicenter, multinational phase Ib/II study (ClinicalTrials.gov identifier: NCT02287233) enrolled patients between January 2015 and May 2017. Data cutoff for efficacy in this analysis was November 8, 2017; cutoff for safety was January 30, 2018. Primary objectives in the dose-escalation phase were to assess safety, pharmacokinetics (PK), maximum tolerated dose, and recommended phase II dose of venetoclax combined with LDAC. In dose expansion, the primary objectives were to obtain preliminary estimates of efficacy: ORR, including CR, CRI, and partial remission (PR); duration of response (DOR); and safety of the combination at the recommended phase II dose. Exploratory objectives were to identify biomarkers of efficacy and resistance.

**Treatment**

Patients were hospitalized and tumor lysis syndrome (TLS) prophylaxis was initiated at least 24 hours before the first dose of venetoclax and continued during a ramp-up period until the target venetoclax dose was reached. Venetoclax was administered orally, once daily, after food. Venetoclax dosing began at 50 or 100 mg and increased over 4 to 5 days to the target venetoclax dose; dosing was continued through day 28 of each cycle. In subsequent 28-day cycles, venetoclax was commenced at the target dose. LDAC (20 mg/m²) was administered by subcutaneous injection once daily, on days 1 to 10. At the completion of a 28-day cycle, if bone marrow blasts were less than 5%, venetoclax dosing was interrupted if needed to promote recovery of neutrophils and platelets. Patients could start the next cycle of therapy when the neutrophil count recovered to least 0.5 x 10⁹/L and the platelet count to at least 25 x 10⁹/L. Once morphologic evidence of leukemia was cleared, granulocyte colony stimulating factor was permitted at investigator’s discretion. Patients could continue receiving treatment until disease progression or until discontinuation criteria were met (Data Supplement).

Because venetoclax is a Cytochrome P450 (CYP3A) substrate, patients receiving CYP3A inhibitors (CYP3A4i) had their venetoclax dose reduced by approximately 50% for moderate CYP3A inhibitors and approximately 75% to 90% for strong inhibitors. If a patient was on multiple inhibitors, venetoclax dose adjustment was based on the strongest CYP3A.

**Study Assessments**

**Safety.** Adverse events (AEs) were graded according to the National Cancer Institute Common Terminology Criteria for Adverse Events Version 4.03. Dose-limiting toxicities (DLTs) were determined during cycle 1 of the dose-escalation phase and defined as grade 4 toxicity, excluding AEs commonly caused by AML (eg, neutropenia, fever). Hematologic DLT was defined as failure of platelet recovery to 25 x 10⁹/L or greater and neutrophils to 0.5 x 10⁹/L or greater within 14 days of the last dose of venetoclax in the absence of residual AML. Clinical and laboratory TLS were considered as previously defined.

**Efficacy.** Bone marrow assessments were performed at screening, after cycles 1 and 3, and then every three cycles until two consecutive samples confirmed CR. Additional bone marrow studies were performed if there was clinical suspicion of recurrence and at final visit. Clinical responses were defined according to International Working Group response criteria for AML (Data Supplement). Pharmacokinetic assessment is described in the Data Supplement. Biomarkers that may be predictive of venetoclax activity and response were assessed during the trial (Data Supplement). Clinical outcome was correlated with the WHO 2008 classification and cytogenetic and molecular markers, including the chromatin-spliceosome, TP53-aneuploidy, and IDH R172 mutation subgroups, as recently proposed. Karyotypic abnormalities were classified using site-reported results, and centrally reported next-generation sequencing data were used to supplement molecular mutation results.
Statistical Analyses
All baseline summary statistics and analyses were based on patient characteristics obtained before initiation of venetoclax or LDAC. The safety population included patients who received at least one dose of venetoclax; the DLT-evaluable population included patients who received at least 80% of planned cycle 1 doses during dose-escalation phase. Further details of statistical analyses are contained in the Data Supplement. For the phase II portion, it was initially planned to enroll 25 patients at the recommended phase II dose to enable a 95% CI for estimation of ORR with margin of error not exceeding plus or minus 25%. The study was later amended to enroll an additional 28 patients to provide further precision regarding the observed response rate estimates, and another 21 patients were enrolled to evaluate patients allowed to receive concomitant strong CYP3A inhibitors if indicated.

RESULTS
All patients enrolled at least 6 months before this analysis. At the time of analysis, the median treatment duration was 42 months (range, 0.2 to 29 months), and the median number of cycles of therapy was five. The median number of cycles of LDAC delivered to patients who achieved CR was seven (range, two to 30), and five (range, one to 16) for patients achieving CRi. Seven patients (9%) remained in the study therapy at time of analysis.

Patient Demographics and Clinical Characteristics
Eighty-two patients were enrolled to a 600 mg venetoclax cohort and received at least one dose of venetoclax. Baseline demographics and clinical characteristics for the 600 mg venetoclax cohort are shown in Table 1. The median age was 74 years (range, 63 to 90 years). Forty-nine percent of patients had secondary AML, and 50% had concomitant CYP3A inhibitor use. Baseline mutations in TP53, FLT3, IDH1/2, and NPM1 were detected in 14%, 23%, 25%, and 13% of patients, respectively.

DLT and Maximum Tolerated Dose
No DLTs were observed in the 600 mg venetoclax cohort during the dose-escalation phase of the study. Of the six patients in the 600 mg dose escalation cohort who proceeded to subsequent cycles, only one patient required dose interruption between cycles 1 and 2, because of thrombocytopenia without residual morphologic AML. At the 800 mg dose level (n = 10), most patients who proceeded to subsequent cycles needed dose interruption between cycles 1 and 2 to permit count recovery, and one patient experienced a hematologic dose-limiting toxicity (grade 4 thrombocytopenia lasting more than 42 days without evidence of residual leukemia in a patient with AML secondary to myeloproliferative neoplasm). Therefore, the recommended phase II dose was determined to be venetoclax 600 mg when combined with LDAC.

Safety Profile
A summary of treatment-emergent AEs for the venetoclax 600 mg cohort is shown in Table 2; further breakdown is available in the Data Supplement. Consistent with expectations for AML, the common grade 3 or 4 AEs, irrespective of cause, were frequently hematologic and included febrile neutropenia (42%), thrombocytopenia (38%), neutropenia (27%), and anemia (27%). The most common nonhematologic AEs of any grade or cause were nausea (70%), diarrhea (49%), hypokalemia (48%), and fatigue (43%). Serious AEs other than AML progression, occurring in at least 5% of patients, were anemia (31%), febrile neutropenia (27%), pneumonia (10%), and sepsis (7%).

Forty-five (55%) patients had venetoclax dose interruptions due to AEs, most commonly between subsequent 28-day cycles (due to delayed neutrophil and platelet recovery in eight and 10 patients, respectively). Dose reductions were necessary in six patients (7%), the majority due to thrombocytopenia. Reduced duration of venetoclax administration to 21 and 14 days occurred in 25 patients (30%) and 14 patients (17%), respectively. Laboratory-defined TLS was reported in two patients (elevations of potassium and phosphorus in one patient and elevations of uric acid and phosphorus in the other), although both were able to complete the venetoclax ramp-up to the intended dose. No clinical TLS was reported. The 30-day mortality rate was 6% (n = 5).

Efficacy
In phase I and II, a total of 82 patients were enrolled to the venetoclax 600 mg dose level. The CR/CRi rate was 54% (95% CI, 42% to 65%); CR was achieved in 26% of patients, and CRi in 28% (Fig 1). Median time to first CR/CRi was 1.4 months (range, 0.8 to 14.9 months). Patients with de novo AML and intermediate-risk cytogenetic features and without prior HMA exposure had the highest rates of CR/CRi: 71%, 63%, and 62%, respectively (Fig 1). A complete breakdown of responses by baseline patient features is shown in the Data Supplement. Among patients achieving CR/CRi after venetoclax plus LDAC, the median duration of remission was 8.1 months (95% CI, 5.3 to 14.9 months; Fig 2A). The median OS for all patients was 10.1 months (95% CI, 5.7 to 14.2 months; Fig 2B). Observed survival for the study population was better for patients achieving CR, CRi, or both (Fig 2C). The 12-month estimated survival rate for those who achieved CR, CRi, or no response was 100%, 49%, 73%, and 5%, respectively (Fig 2C, arrows). Patients without prior HMA exposure had a longer median OS (13.5 months; 95% CI, 7.0 to 18.4 months) than patients with AML previously exposed to prior HMA (4.1 months; 95% CI, 2.9 to 10.1 months; Fig 2D).

Most patients who achieved remission became transfusion independent. Among patients treated with venetoclax in combination with LDAC, 46% (38 of 82) achieved independence from both RBC and platelet transfusion.
Forty-eight percent (39 of 82) achieved RBC transfusion independence, and 60% (49 of 82) achieved platelet transfusion independence. Among the patients who were dependent on transfusions before enrollment, 43% (23 of 53) of those dependent on RBC transfusions and 65% (15 of 23) of those dependent on platelet transfusions became transfusion independent while in the study. The time required for platelet (≥ 50 × 10^9/L) and neutrophil (≥ 0.5 × 10^9/L) recovery was assessed among patients who achieved CR/CRi (Data Supplement). The median time to platelet recovery was 28 days, with 90% or more reaching this threshold by day 53. The median time to absolute neutrophil count recovery was 32 days, with 90% recovered by day 64.

**DISCUSSION**

Optimizing treatment selection for older adults with AML remains challenging. Higher incidence of treatment-related complications is expected from intensive chemotherapy because poor performance status and comorbidities are more prevalent in the elderly. Poor outcomes with intensive chemotherapy are also typical for older patients with poor-risk cytogenetics, monosomal karyotype, secondary AML, or MDS after failure of prior HMA therapy. For patients older than 65 years receiving intensive chemotherapy, reported remission rates range between 45% and 57% and median OS between 5 and 12 months. Low-intensity therapies, such as LDAC monotherapy, have historically been administered with palliative intent, with CR/CRi rates of 11% to 19% and median OS of approximately 5.5 months. Therefore, the observed CR/CRi rate of 54% and 10.1-month median survival for patients treated with venetoclax plus LDAC seems favorable. Among patients older than 75 years, the rate of CR/CRi remained 60%, with median OS of 14.9 months. Importantly, the higher clinical response rate was achieved without an increase in early mortality: the 30-day mortality rate was 6% in the first month after LDAC plus venetoclax treatment, compared with 8% to 13% with LDAC alone from literature reports. This raises the possibility of also evaluating venetoclax combined with low-intensity therapy in younger and/or more fit patients with AML, with the goal of sparing patients the greater toxicity generally observed with intensive chemotherapy.
TABLE 2. Summary of Treatment-Emergent AEs

<table>
<thead>
<tr>
<th>AE with grade ≥ 3</th>
<th>Any AE</th>
<th>Venetoclax 600 mg + LDAC (n = 82)</th>
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</thead>
<tbody>
<tr>
<td></td>
<td>Venetoclax</td>
<td>CR</td>
</tr>
<tr>
<td></td>
<td>82 (100)</td>
<td>63 (78)</td>
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<tr>
<td>Febrile neutropenia</td>
<td>34 (42)</td>
<td>35 (43)</td>
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<tr>
<td>Thrombocytopenia</td>
<td>31 (38)</td>
<td>29 (36)</td>
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<tr>
<td>WBC count decreased</td>
<td>28 (34)</td>
<td>27 (33)</td>
</tr>
<tr>
<td>Anemia</td>
<td>22 (27)</td>
<td>29 (36)</td>
</tr>
<tr>
<td>Neutropenia</td>
<td>22 (27)</td>
<td>33 (41)</td>
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<tr>
<td>Platelet count decreased</td>
<td>20 (24)</td>
<td>28 (34)</td>
</tr>
<tr>
<td>Lymphocyte count decreased</td>
<td>15 (18)</td>
<td>13 (16)</td>
</tr>
<tr>
<td>Neutrophil count decreased</td>
<td>14 (17)</td>
<td>13 (16)</td>
</tr>
<tr>
<td>Hypophosphatemia</td>
<td>13 (16)</td>
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<td>Hypokalemia</td>
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<td>Hypertension</td>
<td>9 (11)</td>
<td>10 (13)</td>
</tr>
<tr>
<td>Pneumonia</td>
<td>9 (11)</td>
<td>11 (14)</td>
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<tr>
<td>Sepsis</td>
<td>9 (11)</td>
<td>6 (8)</td>
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<tr>
<td>Serious AE</td>
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<td>N = 49</td>
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<tr>
<td>Anemia</td>
<td>25 (31)</td>
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<tr>
<td>Febrile neutropenia</td>
<td>22 (27)</td>
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<tr>
<td>Pneumonia</td>
<td>8 (10)</td>
<td></td>
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<tr>
<td>AML progression</td>
<td>7 (9)</td>
<td></td>
</tr>
<tr>
<td>Sepsis</td>
<td>6 (7)</td>
<td></td>
</tr>
</tbody>
</table>

NOTE. Data are presented as No. (%). Adverse events (AEs) were listed if they were reported for at least 10% of patients with grade 3 or higher or at least 5% of patients with serious AEs.

Abbreviation: AML, acute myeloid leukemia.

The outcomes achieved with venetoclax plus LDAC also compare favorably with those obtained during recent studies of HMA monotherapy in older patients with AML: decitabine resulted in CR/CRi of 25.6% and median OS of 7.7 months in older adults with newly diagnosed AML4; azacitidine achieved CR/CRi of 27.8% and median OS of 10.4 months in another phase III study, which included patients either eligible or ineligible for intensive chemotherapy.7 Importantly, both historical studies excluded patients who had received prior HMA for MDS, whereas in the current study, 29% of patients had previously received an HMA. For HMA-naive participants in the current study, the CR/CRi rate was 62%, with a median duration of response 14.8 months and median OS of 13.5 months.

Historical rates of CR/CRi for patients treated with LDAC, azacitidine, or decitabine monotherapy are between 11% and 26%,5,7 with moderate difference in CR rates observed between patients with intermediate-risk versus poor-risk cytogenetics (reported differences between 3% and 13%).8,9 In the current study, rates of CR/CRi were 63% and 42% for patients with intermediate-risk and poor-risk cytogenetics, respectively. Among patients with mutant NPM1, the CR/CRi rate was 89% (eight of nine), which included two patients with coexisting FLT3 and IDH1/2 mutations and three additional patients with either mutant FLT3 or IDH1/2; all eight patients remain alive at more than 1 year. Although larger validation studies are needed, these preliminary observations suggest that such patients may be especially responsive to venetoclax plus LDAC therapy.

Responses to venetoclax monotherapy have been previously reported in patients with IDH1 or IDH2 mutations.46 In the current study, venetoclax plus LDAC resulted in a CR/CRi rate of 72% (13 of 18) and median OS of 19.4 months among patients with IDH1/2-mutant AML. Therefore, venetoclax plus LDAC may also represent a useful treatment strategy for patients with treatment-naive IDH1/2-mutant AML who are ineligible for intensive chemotherapy; further integration of IDH1/2 inhibitors into this regimen for patients with persistent IDH-positive disease may be an objective for future clinical trials. Among patients with TP53 mutation or poor cytogenetic risk, the CR/CRi rate with LDAC and venetoclax was 30% (three of 10) or 42% (11 of 26), respectively, indicating that these historically identified features of poor prognosis remain relevant for this combination.

Venetoclax plus LDAC was associated with rapid achievement of CR/CRi, with median time to first response of 1.4 months, compared with 3.1 months for LDAC alone4 and 3.5 to 4.1 months with HMA therapies.37 However, clinical vigilance and robust supportive care measures are necessary until blast clearance and hematologic recovery are achieved. At the onset of this trial, antifungal prophylaxis with azole antifungals (CYP3A inhibitors) was prohibited because of the potential of increasing venetoclax exposure.41 However, separate pharmacokinetic studies reported that...
venetoclax can be administered, with appropriate dose reductions, to patients with AML using concomitant moderate or strong CYP3A inhibitors (such as the antifungal azoles voriconazole or posaconazole). Consequently, a cohort was added to prospectively evaluate the impact of allowing CYP3A inhibitors when medically required. As such, 50% of the patients enrolled and treated had concomitant CYP3A inhibitor use; no increase in adverse events related to potential CYP3A inhibitor interaction was observed.

Because patients with prior HMA exposure were included in the current study, it is difficult to directly compare efficacy between the current study and historical results. Clinical trials of low-intensity therapies have historically excluded patients who received prior treatment with HMAs, and such patients have an especially poor prognosis. In the current study, nearly one-third (29%) of patients had prior HMA exposure. The CR/Cri rate for those patients with prior HMA exposure was 33%, comparable to rates reported for intensive chemotherapy or CPX-351 for patients experiencing disease progression after prior HMA therapy for antecedent conditions. Among patients without prior HMA treatment, the CR/Cri rate for venetoclax plus LDAC was 62%, which is similar to the reported CR/Cri rate of 67% in a recently published study that combined venetoclax with HMAs for patients with AML. As such, both venetoclax studies, in combination with LDAC or HMAs, support a role for venetoclax as an integral component of standard therapy, especially in older adults with AML. International randomized studies are currently underway for venetoclax or placebo combined with either LDAC (ClinicalTrials.gov identifier: NCT03069352) or azacitidine (ClinicalTrials.gov identifier: NCT02993523).
Venetoclax With LDAC for Older Adults With AML

In conclusion, the combination of venetoclax and LDAC is tolerable and associated with high rates of remission in patients with previously untreated AML who are ineligible for intensive chemotherapy. The high remission rate, low early mortality, rapid induction of remission, and durable length of remission make the combination with venetoclax an attractive and novel treatment option for older adults not suitable for intensive chemotherapy.

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AUTHORS’ DISCLOSURES OF POTENTIAL CONFLICTS OF INTEREST

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Employment: AbbVie/Genentech
Stock and Other Ownership Interests: AbbVie/Genentech
Research Funding: AbbVie/Genentech
Travel, Accommodations, Expenses: AbbVie/Genentech
Other Relationship: AbbVie/Genentech

Rod Humerickhouse
Employment: AbbVie
Stock and Other Ownership Interests: AbbVie

Wan-Jen Hong
Employment: Genentech
Stock and Other Ownership Interests: Roche
Travel, Accommodations, Expenses: Genentech

John Hayslip
Employment: AbbVie
Stock and Other Ownership Interests: AbbVie

Gail J. Roboz
Consulting or Advisory Role: Amphivena Therapeutics, Janssen, Amgen, Astex Pharmaceuticals, Celgene, Genoptix, MedImmune, Novartis, Pfizer, AbbVie, argenx, Array BioPharma, Bayer, Celltrion, Jazz Pharmaceuticals, Orsenix, Genentech, Sandz, Actinium Pharmaceuticals, Astellas Pharma, Eisai, Bayer, Daiichi Sankyo, MEI Pharma, Otsuka Pharmaceutical, Takeda, Roche
Research Funding: AbbVie (Inst), Agios Pharmaceuticals (Inst), Astex Pharmaceuticals (Inst), Celgene (Inst), CTI (Inst), Karyopharm Therapeutics (Inst), MedImmune (Inst), MEI Pharma (Inst), Moffitt (Inst), Novartis (Inst), Onconova Therapeutics (Inst), Pfizer (Inst), Sunesis Pharmaceuticals (Inst), Tensha Therapeutics (Inst), Cellectis (Inst), Janssen (Inst), Amphivena Therapeutics (Inst)
Travel, Accommodations, Expenses: Amphivena Therapeutics, Astex Pharmaceuticals, Janssen, Pfizer, Array BioPharma, Novartis, AbbVie, Jazz Pharmaceuticals, Celgene, Celltrion, Genentech, Sandz, Bayer, Clovis Oncology, Amgen, Sunesis Pharmaceuticals, Eisai

No other potential conflicts of interest were reported.

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