Preclinical Evaluation of the Novel Small-Molecule Integrin α5β1 Inhibitor JSM6427 in Monkey and Rabbit Models of Choroidal Neovascularization

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Objective: To evaluate the pharmacologic activity and tolerability of JSM6427, a potent and first selective small-molecule inhibitor of integrin α5β1, in monkey and rabbit models of choroidal neovascularization (CNV).

Methods: JSM6427 selectivity for α5β1 was evaluated by in vitro binding assays while the ability of JSM6427 to inhibit CNV was investigated in a laser-induced monkey model and a growth factor–induced rabbit model. Intravitreal injections of JSM6427 (100, 300, or 1000 µg) or vehicle were administered immediately after the CNV induction procedure and at weekly intervals for 4 weeks. Fluorescein angiography was performed weekly. Ocular tolerability was evaluated ophthalmoscopically and histologically in both models; additional assessments in monkeys included electroretinography, biomicroscopy, pathological examination, and analysis of JSM6427 pharmacokinetics.

Results: JSM6427 was highly selective for the α5β1–fibronectin interaction. Weekly intravitreal injections of JSM6427 resulted in a statistically significant dose-dependent inhibition of CNV in laser-induced and growth factor–induced models without any ocular JSM6427-related adverse effects. JSM6427 was cleared through the systemic circulation with no evidence of systemic accumulation.

Conclusions: Intravitreal JSM6427 provided dose-dependent inhibition of CNV in monkey and rabbit experimental models.

Clinical Relevance: JSM6427 may provide a new approach for the treatment of ocular neovascular diseases such as age-related macular degeneration in humans.

cells cultured on fibronectin and to inhibit fibronectin-mediated signaling in cultured RPE cells. Systemic administration of JSM6427 induced regression of neovascular lesions in a murine model of CNV, while an α5β1-blocking monoclonal antibody inhibited laser-induced CNV in a monkey model. Together these findings support a role for α5β1 in ocular neovascular diseases such as AMD. This article extends these findings by assessing the specificity of JSM6427 in blocking the α5β1-fibronectin interaction and in evaluating the tolerability and dose response of intravitreally administered JSM6427 in 2 animal models of CNV, including a laser-induced monkey model and a vascular endothelial growth factor (VEGF)/bFGF–induced rabbit model.

COMPETITIVE INTEGRIN BINDING ASSAY

The selectivity of JSM6427 for α5β1 was assessed by means of a previously described competitive binding assay in which soluble integrins (α5β1, αvβ3, αvβ5, α3β1, and α3β1) were incubated with the corresponding bound ligands (fibronectin, vitronectin, fibrinogen, or laminin) in the presence of varying concentrations of JSM6427. The inhibition curves were analyzed using a commercially available software program (SoftMaxPro, version 4.0; Molecular Devices Corp, Sunnyvale, California) to determine the 50% inhibitory concentration (IC50).

LASER-INDUCED CNV IN MONKEYS

Twenty adult cynomolgus monkeys (10 males and 10 females) were assigned to groups by a stratified randomization scheme. For all procedures except ophthalmoscopic examinations (in which only intramuscular ketamine hydrochloride was used), monkeys were sedated with intramuscular ketamine and intravenous ketamine/paracaine in each eye, CNV was induced by laser photocoagulation (Oculight GL 532 with IRIS Medical Portable Slit Lamp Adaptor; Iridex Corp, Mountain View, California). A fundus contact lens was used for the laser treatment. Nine lesions (75 µm, 500-700 mW, 100-200 milliseconds) were placed perimacularly in a grid formation; a vaporization bubble indicated the disruption of Bruch's membrane.

Tobramycin ophthalmic solution was applied topically after photocoagulation. Four groups of 4 monkeys each (2 males and 2 females) received weekly 50-µL intravitreal injections (27-g needle) of JSM6427 (100, 300, or 1000 µg per eye) or vehicle (sterile buffered saline; 130mM sodium chloride, 10mM sodium phosphate buffer, pH 7.2) in each eye. The first dose was given immediately after laser treatment, with additional injections on days 8, 15, and 22. Tobramycin was applied topically immediately after injection and on posttreatment day 2. A fifth group of 4 monkeys served as un.injected controls.

Fluorescein angiography was conducted before the study start and each week thereafter. Fluorescein sodium (10% solution; Alcon, Fort Worth, Texas) was injected via an implanted femoral vein catheter routed to a subcutaneous access port. Photographs were taken with a retinal camera (TRC-50EX; Topcon Medical Systems, Paramus, New Jersey) and captured with an imaging system (IMAGEnet; Topcon) during the arterial phase, the early arteriovenous phase, and several late arteriovenous phases; images were assessed by a masked investigator. Grading for lesion leakage was as follows: 0 (no hyperfluorescence), 1 (mild speckled hyperfluorescence), 2 (moderate hyperfluorescence), 3 (mild lesion leakage), 4 (moderate, clinically significant lesion leakage), or 5 (extensive, clinically significant lesion leakage). The total number of lesions with grade 4 or 5 leakage and the percentage of lesions with grade 4 or 5 leakage in relation to the total were determined.

Direct ophthalmoscopic examinations were performed after each injection (days 1, 8, 15, and 22). Before the study and on days 2, 9, 16, 23, and 30, both slitlamp microscopy and indirect ophthalmoscopy were performed. Eyes were scored by means of a modified Hackett-McDonald scale. Intravascular pressure was measured before the study and approximately 2 hours after injection by pneumotonometer (Model 30 Classic; Medtronic, Jacksonville, Florida) after application of proparacaine.

Electroretinograms (Epic 2000 with Ganzfeld apparatus; LKC Technologies, Gaithersburg, Maryland) were recorded before the study and on days 3 and 31. Monkeys were dark-adapted for 20 to 30 minutes before sedation, and tropicamide and proparacaine were administered topically. Electroretinogram probes were placed subcutaneously beneath each eye and posterior to the brow; contact lenses were placed on the eye with carboxymethylcellulose. Four light sequences (blue, red, white scotopic, and white flicker) were repeated at least 5 times, with the instrument averaging the conductance signals. The electroretinogram recordings were evaluated by a veterinary ophthalmology consultant. All monkeys were killed on study day 32; complete gross and histopathological examinations were performed. Eyes were injected and stored in Davidson fixative and transferred to 10% neutral-buffered formalin within 24 to 48 hours for at least 48 hours and then paraffin embedded. Five step-sections (approximately 30 µm) were stained with hematoxylin-eosin and examined by a veterinary pathologist.

Blood samples were collected before dosing and at 1, 4, 8, 12, and 24 hours after dosing on days 1 and 22. Plasma was evaluated for JSM6427 by a high-performance liquid chromatographic–mass spectrometric assay with a detection limit of 3 ng/mL. Individual plasma concentration–time data were analyzed by noncompartmental pharmacokinetic methods via modeling and analysis software (WinNonline 1.5, Model 200; Pharsight Corp, Mountain View, California) with the use of nominal time points. Areas were calculated by the linear trapezoidal rule.

GROWTH FACTOR–INDUCED CNV IN RABBITS

Choroidal neovascularization was induced by implanting VEGF/bFGF pellets as previously described. Briefly, 15 µg of human recombinant VEGF165 and 15 µg of recombinant bFGF (PeproTech, Rocky Hill, New Jersey) or a phosphate-buffered saline control were incorporated into polymer pellets (Hydroxyn type NCC; HydroMed Sciences, Cranbury, New Jersey). Four groups of 6 adult Dutch-belted rabbits each were anesthetized (intramuscular ketamine hydrochloride/xylazine plus topical tropicamide, phenylephrine hydrochloride, and proparacaine) and VEGF/bFGF pellets were implanted in their right eye, while a fifth group of 6 rabbits received control pellets. Povidone iodine was used for asepsis. The pellets were implanted in the suprachoroidal space. After a conjunctival peritomy in the supratemporal quadrant, a full-thickness sclerotomy (avoiding the choroid) was performed concentric to and 3 mm posterior to the limbus. The implant was placed between the 9- and 10-o’clock positions in an artificial suprachoroidal space created by passing a cyclodialysis spatula between the choroid and sclera. Immediately after implantation of the active pellets and on days 7, 14, and 21, the 4 groups of rabbits received in the same eye a midvitreous 50-µL intravit-
real injection (30-g needle, between the 1- and 2-o’clock positions) of JSM6427 (100, 300, or 1000 µg per eye) or vehicle alone followed by topical 0.3% ciprofloxacin (Alcon).

Rabbits were examined by indirect ophthalmoscopy preoperatively, postoperatively, and weekly thereafter for 4 weeks; observations were graded by the McDonald and Shadduck system.28 Standard color fundus photography and fluorescein angiograms using a retinal camera (TRC 50EX; Topcon Medical Systems) were performed before the study and weekly thereafter. Fluorescein sodium (10% solution) was administered intravenously through the ear vein, and photographs were taken during a 30-second period for up to 10 minutes. Grading for lesion leakage was as follows: (1) total area of the lesion; (2) severity of leakage/exudation over time on a scale of 0 (no leakage), 1 (mild), 2 (moderate), and 3 (robust); and (3) a binary assessment of CNV presence (0, no; 1, yes). Histopathology of formalin-fixed tissues was performed.

ANIMAL USE STATEMENT

All procedures were conducted in accord with the Association for Research in Vision and Ophthalmology Statement for the Use of Animals in Ophthalmic and Vision Research and the US Department of Agriculture Animal Welfare Act as well as conditions specified in the Guide for the Care and Use of Laboratory Animals (1996). The protocol was approved by the facility institutional animal care and use committees (North American Science Associates Inc, Irvine, California, and Charles River Laboratories Preclinical Services, Sparks, Nevada).

STATISTICAL ANALYSIS

Statistical differences between groups were determined by 1-way analysis of variance. If significant, analysis of variance was followed by the Newman-Keuls multiple comparison test using statistical software (Prism 4; GraphPad Software, San Diego, California). All lesions (grades 1-5) were included in the calculation, with P < .05 considered statistically significant.

RESULTS

JSM6427 COMPETITIVE BINDING ASSAY

JSM6427 showed highly selective inhibition of the α5β1-fibronectin interaction in vitro, with an IC50 of 0.57 nmol/L (Table). JSM6427 was at least 1700-fold less potent in inhibiting the binding of other integrins to their corresponding ligands.

Table. The IC50 of JSM6427 for Each of the Integrin-Ligand Pairs

<table>
<thead>
<tr>
<th>Integrin</th>
<th>Concentration, Mean (SD), nmol/L</th>
<th>Ligand</th>
<th>Selectivity Factora</th>
</tr>
</thead>
<tbody>
<tr>
<td>α5β1</td>
<td>0.57 (0.35)</td>
<td>Fibronectin</td>
<td>NA</td>
</tr>
<tr>
<td>αvβ3</td>
<td>980 (700)</td>
<td>Vitronectin</td>
<td>1700</td>
</tr>
<tr>
<td>αvβ5</td>
<td>4300 (3900)</td>
<td>Vitronectin</td>
<td>7500</td>
</tr>
<tr>
<td>α1β8β3</td>
<td>&gt;50 000</td>
<td>Fibronogen</td>
<td>&gt;50 000</td>
</tr>
<tr>
<td>α3β1</td>
<td>53 000</td>
<td>Laminin</td>
<td>~100 000</td>
</tr>
</tbody>
</table>

Abbreviations: IC50, 50% inhibitory concentration; NA, not applicable.

aSelectivity was calculated according to the following formula: IC50 for other integrin-ligand combinations/IC50 of JSM6427 for α5β1-fibronectin.

JSM6427 INHIBITION OF LASER-INDUCED CNV IN MONKEYS

By week 2 after photocoagulation, clinically significant (grade 4/5) leakage developed in all groups (Figure 1A). While the number of grade 4/5 lesions in the untreated control group was identical to that in the vehicle-injected control group (30 each), there were reduced numbers of lesions in all JSM6427-treated groups by week 2 (n = 16, n = 14, and n = 7 for the 100-, 300-, and 1000-µg groups, respectively). There were similar results at weeks 3 and 4 (data not shown). Representative fluorescein angiographic photographs of eyes treated with vehicle or 1000 µg of JSM6427 during the 4-week time course are shown in Figure 2.

By week 2, the percentage of clinically significant lesions in JSM6427-treated eyes in relation to the total number of lesions was lower than in uninjected control or vehicle-injected eyes (42% for both untreated and...
vehicle-treated controls and 25%, 19%, and 10% for the 100-, 300-, and 1000-µg groups, respectively. These effects were maintained through weeks 3 (Figure 1B) and 4 and were statistically significant for the high-dose group compared with the vehicle-treated group (P < .05) for all measured time points.

OCULAR AND SYSTEMIC TOXIC EFFECTS IN MONKEYS

No ophthalmologic findings were attributed directly to JSM6427 administration. Findings attributed to the injection procedure were posterior subcapsular cataract (1 of 8 eyes in both the 100- and 300-µg treatment groups and 3 of 8 in the vehicle group) and severe inflammation (2 of 8 in the 100-µg treatment group after the second and third injections). Two eyes receiving 1000 µg of JSM6427 developed transient vitreous haze (mild in 1 and moderate in 1) that resolved by day 9. Mild conjunctival congestion/discharge, white opacities resembling cells on the posterior lens capsule, and mild hemorrhage associated with laser burns also occurred across treatment groups. Intraocular pressure levels were maintained within normal limits (15-30 mm Hg).

No abnormalities in electroretinogram responses attributable to intravitreal JSM6427 were noted (data not shown). At day 31, individual eyes in the vehicle-treated group exhibited decreases in amplitudes greater than 25% of the baseline values, which were attributed to individual variability and the injection procedure. Abnormal responses at day 31 in 2 eyes in the 100-µg group were considered secondary to procedure-related inflammation.

There were no gross or microscopic pathological ocular findings related to the administration of JSM6427. Focal/multifocal laser-induced lesions were as expected after retinal laser photocoagulation. Histologic findings believed related to laser-induced retinal and choroidal lesions included small numbers of erythrocytes in the vitreous or optic nerves, and findings attributable to the injection procedure included minimal to mild fibroplasia/fibrosis of the ciliary body or choroid. Incidental lesions included minimal to mild, focal/multifocal infiltrates of mixed mononuclear inflammatory cells in the ciliary body, choroid, sclera, cornea, and conjunctiva of numerous animals, including the controls. Injection-related unilateral panophthalmitis was observed in 2 of 8 eyes in the 100-µg treatment group. There were no JSM6427-related nonocular clinical, toxicologic, or histopathological findings.

PHARMACOKINETICS OF JSM6427 AFTER INTRAVITREAL ADMINISTRATION IN MONKEYS

Weekly intravitreal injections of JSM6427 resulted in very low plasma levels on days 1 and 22. JSM6427 was detected consistently only in plasma samples of animals given 300 or 1000 µg per eye. After administration of JSM6427 at 300 and 1000 µg per eye per dose, mean plasma concentrations reached Cmax at a mean time of 6 or 8 hours on both days 1 and 22 and then declined. The mean time through which concentrations were measurable on each day was 12 hours at 300 µg per eye per dose and 24 hours at 1000 µg per eye per dose. The mean terminal plasma half-life after dosing of 1000 µg per eye per dose, estimated for a limited set of animals, was approximately 15 hours on days 1 and 22. The mean plasma Cmax on days 1 and 22 ranged between 10.6 and 11.7 ng/mL and between 9.1 and 12.1 ng/mL, respectively, for 300 µg per eye per dose and between 18.4 and 26.6 ng/mL and between 20.3 and 27.4 ng/mL, respectively, for 1000 µg per eye per dose (Figure 3). Mean Cmax (Figure 3) and mean area under the plasma concentration–time curve from time zero to the time of the last measurable concentration (AUC(last)) (data not shown) were slightly different between males and females. Mean Cmax increases were slightly less than the relative increase in dose, while the increases in mean AUC(last) were approximately proportional (for males) or slightly greater (for females) than the relative 3.3-fold increase in dose.

Mean Cmax and mean AUC(last) values on day 22 were comparable with values on day 1, indicating the absence of accumulation in this study. Earlier studies (data not shown) demonstrated a very rapid elimination of JSM6427 after administration.
intravenous administration in the monkey (half-life, 9 minutes), while JSM6427 exhibited flip-flop kinetics after intravitreal injection, with the rate of distribution from the eye being slower than systemic elimination.

**Intravitreal JSM6427 Inhibits Growth Factor–Induced CNV in Rabbits**

In the growth factor–induced rabbit model, which produces stable CNV lesions in which edema is distinct from CNV leakage,29 eyes implanted with VEGF/bFGF-containing pellets developed florid CNV and leakage in vehicle-treated eyes after 2 weeks, increasing through week 3 (Figure 4A and Figure 5). Eyes with control pellets showed no leakage on fluorescein angiography (Figure 4A and Figure 5). At week 3 (Figure 4B), there was a dose-dependent suppression of choroidal vascular leakage in eyes treated with JSM6427 (*P < .05 for control vs 100-, 300-, and 1000-µg groups [Figure 5], respectively). Week 4 differences also were significant (data not shown).

Ophthalmologic and histopathological examinations showed no evidence of JSM6427-related toxic effects. Sporadic minimal or mild focal histologic findings, including retinal disorganization, keratitis, and the presence of macrophages within the retina or choroid, were seen across all treatment groups and were attributed to the injection or implantation procedures.

**Comment**

JSM6427 (or a related molecule) was previously shown to inhibit neovascularization in murine CNV,30 hypoxia-induced,30 and corneal31,32 models, suggesting that the α5β1-fibronectin interaction is an active contributor to angiogenesis in ocular neovascular disease. The present study extends these findings by demonstrating that JSM6427 blocks this interaction with high selectivity and potency, providing a dose-dependent reduction in CNV without evidence of ocular or retinal toxic effects; the efficacy at the highest dose was comparable with that of an α5β1-blocking monoclonal antibody.22 These findings are consistent with mechanistic studies showing that JSM6427 inhibited the proliferation,32 migration,30,32 and sprouting30,32 of endothelial cells in vitro and induced apoptosis of endothelial cells in vitro and in CNV tissues.38

In neovascular AMD, clinical studies have identified RPE cells in subretinal membranes outside the normal RPE layer.33-35 These activated “transdifferentiated” RPE cells may contribute to the development of CNV by generation of angiogenic and inflammatory growth factors.35 Previous work has shown that α5β1 is highly ex-

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**Figure 4.** Average leakage grade during weekly intravitreal injections of JSM6427 (100, 300, or 1000 µg) or vehicle administered to rabbit eyes after implantation of vascular endothelial growth factor/basic fibroblast growth factor pellets or control pellets without growth factor. A, Average leakage grade for each of the dose groups at each weekly evaluation. B, Average leakage grade at week 3 for each of the dose groups. *P < .05.

**Figure 5.** Representative fluorescein angiograms of rabbit eyes treated with control pellets without growth factor and no injection (A), vascular endothelial growth factor/basic fibroblast growth factor (VEGF/bFGF) pellets and vehicle injections (B), or VEGF/bFGF pellets and 1000-µg JSM6427 injections (C).
pressed on proliferating and migrating, but not quiescent, RPE cells, and that JSME427 was able to inhibit the migration of ARPE-19 cells (unpublished data) and fetal RPE cells. Moreover, expression of the a5 subunit was found to be significantly upregulated on RPE cells by tumor necrosis factor a, which has been found to be highly expressed on CNV membranes. By inhibiting RPE proliferation and migration, JSME427 might have additional effects on the growth of subretinal membranes independent of its effects on endothelial cells.

In addition to being expressed on proliferating RPE cells, a5beta1 expression is upregulated on activated mononuclear cells, which are believed to play a key role in CNV by producing growth factors and cytokines, and on myofibroblasts. Therefore, both anti-inflammatory and anti-fibrotic effects of JSME427 may have contributed to the inhibition of neovascularization by JSME427 in our laser model, which approximates CNV related to AMD. In AMD, fibrosis can lead to destruction of the sensory retina, RPE, and inner chorioid with the formation of a fibrovascular disciform scar, resulting in profound vision loss. Further, ocular neovascular diseases such as AMD are suspected to have an inflammatory component, and on myofibroblasts. Therefore, both anti-inflammatory and anti-fibrotic properties of JSME427 warrant further investigation.

A notable function of integrins is that they provide a linkage between extracellular molecules and the actin-based microfilament system; in many cases, proliferative responses to soluble growth factors are dependent on integrin-mediated adherence to extracellular substrates. Blocking angiogenesis through inhibition of integrin-mediated signaling has the potential to inhibit the cellular responses to growth factors as well as cytokines and other inflammatory mediators. The findings of the present study demonstrating that JSME427 reduced the formation of CNV induced by sustained suprachoroidal release of VEGF and bFGF support a new mode of action for JSME427 that differentiates it from recent therapeutic approaches based on VEGF inhibition.

The results of the pharmacokinetic analysis showed that there was detectable systemic exposure after intravitreal injection of JSME427 in monkeys, rapidly declining over 24 hours. For application in clinical trials, it is expected that there will be limited systemic effects because of the very short systemic exposure to low concentrations of JSME427. In addition, a5beta1 integrin expression in normal tissues is very limited, suggesting that the anti-inflammatory and anti-fibrotic properties of JSME427 warrant further investigation.

Subsequent to the positive results of these preclinical studies, a phase 1 clinical trial evaluating intravitreal JSME427 in treating neovascular AMD is under way (clinicaltrials.gov identifier: NCT00536016). The trial includes a single-dose study (ranging from 0.15-1.5 mg per eye) and a repeated-dose study in which 4 weekly injections with the same dose range will be administered. In addition, due to its mechanism of action, JSME427 may show promise not only as a treatment for AMD but also for other ocular neovascular diseases such as proliferative diabetic retinopathy and proliferative vitreoretinopathy.

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Author Contributions: Dr Zahn had full access to all the data in the study and takes responsibility for the integrity of the data and the accuracy of the data analysis.

Financial Disclosure: Dr Zahn is an employee of Jerini AG and is a patent designee; Dr Vossmeyer is an employee of Jerini AG; Dr Stragies is an employee of Jerini AG and is a patent designee; Ms Wills is a consultant for Jerini AG; Dr Wong is a consultant for Jerini AG; Dr Löffler received financial support from Jerini AG; Dr Adams is an employee of Jerini Ophthalmic Inc; Dr Knolle is a former employee of Jerini AG and is a patent desigee.

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