Rapid and Sustained Long-Term Efficacy and Safety of Canakinumab in Patients With Cryopyrin-Associated Periodic Syndrome Ages Five Years and Younger

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Objective. To assess long-term efficacy and safety of canakinumab and the response to vaccination in children ages ≤5 years with cryopyrin-associated periodic syndrome (CAPS).

Methods. CAPS patients (ages ≤5 years) received 2 mg/kg canakinumab subcutaneously every 8 weeks; patients with neonatal-onset multisystem inflammatory disease (NOMID) received a starting dose of 4 mg/kg in this open-label trial. Efficacy was evaluated using physician global assessment of disease activity and serum levels of C-reactive protein (CRP) and amyloid A (SAA). Adverse events (AEs) were recorded. Vaccination response was evaluated using postvaccination antibody titers at 4 and 8 weeks after immunization.

Results. Of the 17 patients enrolled, 12 (71%) had Muckle-Wells syndrome, 4 (24%) had NOMID, and 1 (6%) had familial cold autoinflammatory syndrome. All 17 patients had a complete response to canakinumab. Disease activity improved according to the physician global assessment, and for 65% of the patients autoinflammatory disease was characterized as “absent” at the end of the study. Median CRP levels decreased over time. No such change was evident in SAA levels. During the extension study, postvaccination antibody titers increased above protective levels in 16 (94%) of 17 assessable vaccinations. Ten of the patients (59%) had AEs suspected to be related to canakinumab; 8 (47%) experienced at least 1 serious AE (SAE). None of the AEs or SAEs required interruption of canakinumab therapy.

Conclusion. Our findings indicate that canakinumab effectively maintains efficacy through 152 weeks and appears to have no effect on the ability to produce antibodies against standard childhood non-live vaccines. The safety profile of canakinumab was consistent with previous studies, supporting long-term use of canakinumab for CAPS in children ≤5 years of age.
INTRODUCTION

Cryopyrin-associated periodic syndrome (CAPS) is a rare autosomal-dominant autoinflammatory disorder that includes a group of 3 overlapping inflammatory disorders: familial cold autoinflammatory syndrome (FCAS), Muckle-Wells syndrome (MWS), and chronic infantile neurologic, cutaneous, articular (CINCA) syndrome/neonatal-onset multisystem inflammatory disease (NOMID) (hereafter referred to as NOMID) (1). CAPS is caused by gain-of-function mutations in the NLRP3 gene, which lead to increased production of interleukin-1β (IL-1β) (2–4). The clinical manifestations of CAPS include characteristic urticaria-like rash, recurrent fever episodes, ocular inflammation, musculoskeletal manifestations, and central nervous system (CNS) involvement (5). The milder symptoms include repeated episodes of fever and chills with malaise, intense fatigue, headaches, conjunctivitis, diffuse urticaria-like rash, and generalized limb pain (6). These symptoms, when considered separately, often lead to a diagnostic delay of CAPS, which compromises quality of life and exposes patients to the risk of neurologic complications (including deafness) and, in the longer term, renal failure from secondary AA amyloidosis (7–9).

Early diagnosis of autoinflammatory diseases and effective treatment to control inflammation and prevent irreversible damage, such as deafness, severe joint deformity, growth retardation, and AA amyloidosis, are critical (10). With the well-established efficacy of IL-1 blockade and the widespread availability of genetic testing (helpful in atypical or complex cases), patients with CAPS are increasingly being diagnosed and treated in early life (5). Although the later-stage manifestations related to NOMID, such as bony overgrowth and bone deformities, are not reversible, most CAPS-specific symptoms may be reversible if patients are treated early. Growth retardation, CNS inflammation, and hearing loss have been reported to improve in some patients if treatment is provided in a timely manner (10–12). Early detection and consequent treatment for CAPS patients are critical to improve the patient’s quality of life.

Current data on treatment in infants and children <2 years of age are limited and largely based on case reports (13,14). Prior to the widespread use of biologic agents, basic treatments to relieve canakinumab subcutaneously every 8 weeks for the entire study period (including the core and extension studies). Patients in whom a complete response was not achieved after the first dose of canakinumab, and those who experienced a relapse before the next scheduled dose, were eligible for a stepwise dose titration (4 to 6 to 8 mg/kg). For patients previously treated with an anti–IL-1 agent and for NOMID patients, a starting dose of 4 mg/kg was administered to mitigate against the risk of rebound flares anecdotally observed following withdrawal of IL-1 blockade in CAPS patients, and since more severe CAPS phenotypes have previously been observed to require higher doses of IL-1 blockade, respectively (22,26). The study protocol was reviewed by the independent ethics committee or institutional review board of each center, and the study was conducted according to the ethics principles of the Declaration of Helsinki. Written informed consent was obtained from each patient’s legal guardian before randomization.

Patients. Inclusion criteria. Patients with CAPS ages 28 days to 60 months, with confirmed NLRP3 mutations and a body weight of ≥2.5 kg, were included in the core study. If patients were receiving the anti–IL-1 agents anakinra or rilonacept before enrollment, these were discontinued prior to the baseline visit and the patients had to demonstrate active disease prior to enrollment. Patients who completed the core study with no major protocol deviations and were ≥1 year of age were rolled over to the extension study.
**Exclusion criteria.** Preterm neonates, patients with a history of recurrent infections and/or evidence of active infections or latent tuberculosis infection, patients with neutropenia, and those who had been treated with immunosuppressive drugs or received live vaccinations up to 3 months before screening were excluded.

**Assessment of treatment response.** The main objective of the core study was to assess the efficacy of canakinumab with respect to treatment response (Table 1). The extension study assessed the long-term efficacy of canakinumab with respect to relapse in patients who completed the core study. Complete response was defined as a physician global assessment of auto-inflammatory disease activity as less than or equal to “minimal” (on a 5-point scale of absent, minimal, mild, moderate, and severe) and assessment of skin disease as less than or equal to “minimal” (on a 5-point scale of absent, minimal, mild, moderate, and severe) and serologic response indicated by serum C-reactive protein (CRP) <15 mg/liter or serum amyloid A protein (SAA) <10 mg/liter (21). Relapse was defined for complete responders based on clinical features (physician global assessment greater than “minimal” or physician global assessment greater than or equal to “minimal” and assessment of skin disease greater than “minimal”) and serologic features (serum CRP >30 mg/liter or SAA >30 mg/liter).

**Physician global assessment of autoinflammatory disease activity.** For the physician global assessment, the following 8 features were each graded on a 5-point scale (ranging from absent to severe): skin disease (urticaria-like skin rash), arthralgia, myalgia, headache/migraine, conjunctivitis, fatigue/malaise, other symptoms related to autoinflammatory syndrome, and symptoms not related to autoinflammatory syndrome.

**Evaluation of markers of inflammation.** Serum levels of the inflammation markers CRP and SAA were determined in a central laboratory for all patients, regardless of age. The normal ranges for CRP and SAA levels were 0–6 mg/liter and 0–6.7 mg/liter, respectively.

**Safety assessment.** Safety of canakinumab was assessed in terms of adverse events (AEs) and serious AEs (SAEs) according to Medical Dictionary for Regulatory Activities (MedDRA; version 17.1) reporting criteria for clinical trials.

**Immunogenicity assessment.** Anticanakinumab antibodies concentrations were assessed in serum, and their potential correlation with any AEs or SAEs and/or loss of efficacy was analyzed.

**Assessment of antibody titers against vaccine antigen.** The ability to attain or maintain protective antibody levels was assessed for inactivated vaccines, which the patients received as part of national vaccination programs, and thus potentially included the following antigens for the core study: *Corynebacterium diphtheriae*, *Bordetella pertussis*, *Neisseria meningitidis*, *Clostridium tetani*, influenza A (H1N1 and H3N2), influenza B, *Haemophilus influenzae B*, *Streptococcus pneumoniae*, and hepatitis B. The extension study included assessment of antibody titers against diphtheria, pertussis, *Meningococcus*, tetanus, *Haemophilus influenzae B* (polysaccharide or conjugate), influenza, *Streptococcus pneumoniae*, and hepatitis A and B. No live vaccinations were allowed throughout the course of the study or up to 3 months after the last canakinumab dose.

**Assessment of vaccination response.** Patients were assessed for vaccination response if antibody titer was measured 0–14 days after vaccination (referred to as the “predose assessment”), and on at least 1 subsequent occasion (at 4 weeks and/or 8 weeks after vaccination). Patients were not assessed for a vaccination response if the antibody titer was already sufficient before dosing and maintained during the study.

**Assessment of neurologic, ophthalmologic, and auditory features.** At screening and week 24, magnetic resonance imaging (MRI) of the brain and inner ear was performed, and neurologic, ophthalmologic, and auditory brainstem responses were assessed. A final assessment was performed for all evaluations at the end-of-study visits in the core and extension studies.

**Evaluation of pharmacokinetics/pharmacodynamics.** Canakinumab concentration data were collected and analyzed. A pharmacokinetics-binding model was applied to estimate pharmacokinetics parameters such as clearance, apparent volume of distribution, and first-order absorption rate constant for canakinumab in CAPS patients.

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**Table 1.** Assessment of response to canakinumab in patients with cryopyrin-associated periodic syndrome*

| Definition of complete response | PGA of autoinflammatory disease activity less than or equal to “minimal” AND Skin assessment score less than or equal to “minimal” AND CRP <15 mg/liter OR SAA <10 mg/liter |
| Definition of relapse | PGA of autoinflammatory disease activity greater than “minimal” OR PGA of autoinflammatory disease activity greater than or equal to “minimal” and skin assessment score greater than “minimal” AND Serum CRP AND/OR SAA >30 mg/liter |

* Physician global assessment (PGA) of autoinflammatory disease activity and skin assessment scores were summarized by severity code (absent, minimal, mild, moderate, severe). CRP = C-reactive protein; SAA = serum amyloid A.† In patients in whom a complete response had been achieved.
Table 2. Baseline characteristics of the 17 patients with cryopyrin-associated periodic syndrome treated with canakinumab*

| Age, median (range) months | 31 (1–59) |
|---------------------------|-----------|
| Sex, no. (%) male         | 12 (71)   |
| Race, no. (%)             |           |
| Caucasian                 | 16 (94)   |
| Asian                     | 1 (6)     |
| Weight, no. (%)           |           |
| <15 kg                    | 14 (82)   |
| ≥15 kg                    | 3 (18)    |
| Molecular diagnosis of NLRP3 mutation, no. (%) | 17 (100) |

| Phenotype, no. (%)         |           |
| FCAS                      | 1 (6)     |
| MWS                       | 12 (71)   |
| NOMID                     | 4 (24)    |
| Time from diagnosis to study entry, mean ± SD years | 2.6 ± 1.5 |
| Serum markers of inflammation, median (range)† | CRP, mg/liter‡ | 7 (0–165) |
| SAA, mg/liter§            | 9 (0–861) |

* FCAS = familial cold autoinflammatory syndrome; MWS = Muckle-Wells syndrome; NOMID = neonatal-onset multisystem inflammatory disease; CRP = C-reactive protein; SAA = serum amyloid A.
† Data were available for 16 patients.
‡ Normal range 0–6 mg/liter.
§ Normal range 0–6.7 mg/liter.

RESULTS

Patient disposition and baseline clinical characteristics. Seventeen patients (with a median age of 31 months [range 1–59 months]) with CAPS were enrolled in the core study. Six of the patients were younger than 2 years of age. All enrolled patients completed the core study and entered the extension study. Most patients (14 of 17 [82%]) completed the extension study; of the 3 patients who discontinued during the extension study, 2 discontinued owing to unsatisfactory therapeutic effect (and were thus managed outside the clinical protocol as per local standard of care) and 1 had no reason for discontinuation recorded (and data were also subsequently unavailable). Screening for NLRP3 mutations was performed by routine genetic sequencing as part of regular clinical care for all patients. The mean time from diagnosis to study entry was 2.6 years. All 17 patients had at least 1 confirmed pathologic NLRP3 mutation (Supplementary Table 1, available on the Arthritis & Rheumatology web site at http://onlinelibrary.wiley.com/doi/10.1002/art.41004/abstract). An additional TNFRS1 variant (p.R121Q, also referred to as the R92Q variant) was observed in 1 MWS patient. Of the 17 patients enrolled, 12 (71%) had MWS, 4 (24%) had NOMID (based on neurologic involvement in 4, on papilledema in 3, and overall severity of the phenotype in all 4 as judged by the local investigator), and 1 (6%) had FCAS (Table 2). Median baseline CRP and SAA levels were abnormal (7.0 mg/liter for CRP and 9.1 mg/liter for SAA).

Dosage. Patients with the FCAS or MWS phenotype received a starting dose of 2 mg/kg of canakinumab every 8 weeks, with the exception of 3 MWS patients who received a starting dose of 4 mg/kg due to prior exposure to anti-IL-1 agents. Patients with NOMID received a starting dose of 4 mg/kg due to clinical concern about severe phenotype, with the excep-

Canakinumab dose

![Canakinumab dose diagram](image_url)

Figure 1. Canakinumab dosing at baseline, the end of the core study, and the end of the extension study (final dose) in patients with familial cold autoinflammatory syndrome (FCAS), Muckle-Wells syndrome (MWS), and neonatal-onset multisystem inflammatory disease (NOMID). Doses are rounded up to the nearest whole number. Values in the >4 mg/kg category at the end of the core study and for the final dose are the median (range). See Supplementary Table 1, available on the Arthritis & Rheumatology web site at http://onlinelibrary.wiley.com/doi/10.1002/art.41004/abstract for exact doses in individual patients.
tion of 1 patient who received a starting dose of 6 mg/kg as an a priori agreed protocol deviation, again based on clinical concern regarding particularly severe phenotype (Supplementary Table 1, available on the Arthritis & Rheumatology web site at http://onlinelibrary.wiley.com/doi/10.1002/art.41004/abstract). A total of 9 (53%) of 17 patients did not require any dose up-titration during the core study. Three of 4 NOMID patients and 5 of 12 MWS patients required dose up-titration. During the extension study, 7 (41%) of the 17 patients did not require any dose up-titration, whereas 10 patients (6 with MWS, 3 with NOMID, and 1 with FCAS) did require dose up-titration (Figure 1).

**Pharmacokinetics/pharmacodynamics.** Data relating to pharmacokinetics and pharmacodynamics evaluation were analyzed and will be reported separately. Comparison of dose-normalized drug concentrations among patients who received a consistent canakinumab dose demonstrated that exposure to canakinumab in patients ages 2 years or younger (n = 5) was similar to that observed in patients older than 2 years (n = 4).

**Efficacy.** Treatment responses. Overall, a complete response was achieved in 16 (94%) of the 17 patients by the end of the core study (week 56). In the extension study, a complete response was achieved in 100% of the patients within 72 weeks from the start of the study (Figure 2A), and no new relapses occurred after week 96 (Figure 2B).

**Physician global assessment of disease activity.** The physician global assessment of autoinflammatory disease activity in the overall population decreased from baseline to the end of the study (Figure 3). The proportion of patients without autoinflammatory disease (absent) increased from 24% (4 of 17 patients) at baseline in the core study to 65% (11 of 17 patients) at the end of the study. The proportion of patients with mild and moderate autoinflammatory disease decreased from 47% (8 of 17 patients) at baseline to 6% (1 of 17 patients) at the end of the study and from 24% (4 of 17 patients) to 0% at the end of the study, respectively. There was 1 report of severe autoinflammatory disease flare at week 72, but none from week 80 to the end of the study.

**Assessment of skin disease.** The proportion of patients without skin disease (absent) increased from 29% (5 of 17 patients) at baseline in the core study to 94% (16 of 17 patients) at the end of the study. The proportion of patients with moderate skin disease decreased from 29% (5 of 17 patients) at baseline to 0% at the end of the study. Severe skin disease was reported for 6% (1 of 17 patients) at baseline and was not reported for any patient at any visit thereafter.

**Change in levels of serum markers of inflammation (CRP and SAA).** The median CRP level (normal range 0–6 mg/liter) decreased from 7.0 mg/liter at baseline (precanakinumab) (range 0.0–165.0) to 1.0 mg/liter at the end of the core study (range 0.0–22.0) and 3.0 mg/liter at the end of the study (range 0.0–47.0). The median SAA level (normal range 0–6.7 mg/liter) decreased from 9.1 mg/liter at baseline (range 0.0–861.0) to 2.2 mg/liter at the end of the core study (range 0.0–331.0). However, the median SAA level increased to 9.0 mg/liter at the end of the study (range 0.0–173.0).

**Immunogenicity.** Antibodies against canakinumab were not detected in any of the patients in the core or extension studies.

**Findings of vaccination assessments.** In the core study, 7 (41%) of the 17 patients (ages 5–59 months) received 10 types of inactivated vaccines. These 7 patients received a total of 31 vaccinations, 18 of which were assessable for a vaccination response. For the remaining 13 vaccinations, no predose antibody titer was available and, therefore, response was not assessed. All assessable patient vaccination cases showed a positive response (antibody titers increased above protective level) (Supplementary Table 2, available on the Arthritis & Rheumatology web site at http://onlinelibrary.wiley.com/doi/10.1002/art.41004/abstract). For all 31 patient vaccinations, including those without a predose antibody titer, a
protective antibody titer level was observed during the core study, which was maintained throughout the trial. In the extension study, 4 (24%) of the 17 patients (ages 0–5 years) received 8 types of inactivated vaccines. These 4 patients received a total of 20 vaccinations, and 17 of these unique patient vaccinations were assessable for response; 16 (94%) showed a positive response, with antibody titers increasing above the protective level (Supplementary Table 2, available at http://onlinelibrary.wiley.com/doi/10.1002/art.41004/abstract). Furthermore, in 19 patient vaccination cases, including those without a predose antibody titer, a protective antibody titer level was observed during the study, which was maintained throughout the extension study, as seen in the core study. Predose antibody titers and the last measured antibody titer for each patient and corresponding vaccinations during the extension study are shown in Supplementary Table 2.

In one patient who received a Tetravac formulation (diphtheria, tetanus, acellular pertussis, and inactivated polio combination), a positive response to *Bordetella pertussis* and *Corynebacterium diphtheriae* was achieved, but not to *Clostridium tetani*; the values provided from the laboratory represented optical density rather than antibody concentrations and hence were considered nonevaluable. This patient had no AEs associated with this vaccination, nor lymphopenia or neutropenia at the time of last vaccination and antibody titer assessments. There was no disease flare induced by vaccination.

**Neurologic, ophthalmologic, and auditory findings.** Three patients had clinically significant audiology abnormalities at baseline and were also classified as having abnormalities during the extension study. The hearing loss reported for 1 NOMID patient continued to worsen during the extension study.

On neurologic assessment, 4 patients were found to have the following abnormalities: delay in motor function and coordination and language delay in 1 NOMID patient, gap of acquisitions in 1 MWS patient, intention tremors in 1 MWS patient, and lack of responsiveness in 1 MWS patient. The ophthalmologic assessment showed abnormal findings in 2 NOMID patients who had clinically significant abnormalities during the extension study. One patient had uveitis in the right eye, and the second patient had a swollen and thick-ended retinal fiber layer in the left eye, which was recorded as a clinically insignificant abnormality on days 946 and 1,125.

All but 1 patient underwent brain MRI. The findings were interpreted as normal for all of the patients, except for 1 patient with NOMID who had dilatation of ventricles, bilateral T2 hyper-signal in parietal white matter, and delay of myelination on day 579 of the extension study. (At a later assessment, on day 629, this finding was recorded as a clinically insignificant abnormality.)

**Safety.** Overall, the mean exposure was 951 days per patient with a total of 44 patient-years for the core and extension study periods, which comprised a mean exposure of 429 days per patient and 20 patient-years for the core study and 546 days per patient and 25 patient-years for the extension study. In the core study, all 17 patients exposed to canakinumab experienced at least 1 AE. The most common AEs were nasopharyngitis (in 7 [41%] of the 17 patients) and upper respiratory tract infection (in 7 [41%]) (Table 3). Overall,
Table 3. AEs and SAEs experienced by patients with CAPS treated with canakinumab in the core and extension studies*

|                          | Core study | Extension study |
|--------------------------|------------|-----------------|
| Exposure, mean ± SD days† | 428.6 ± 46 | 546.1 ± 210     |
| Total AEs, no. (%)       | 17 (100)   | 16 (94)         |
| Total SAEs, no. (%)      | 4 (2.3)    | 8 (47)          |
| Common AEs, no. (%)‡     |            |                 |
| Naiopharyngitis          | 7 (41)     | 7 (41)          |
| Upper respiratory tract infection | 7 (41) | 0 (0)          |
| Diarrhea                 | 0 (0)      | 7 (41)          |
| Pyrexia                  | 6 (35)     | 6 (35)          |
| Rhinitis                 | 6 (35)     | 0 (0)           |
| Vomiting                 | 0 (0)      | 6 (35)          |
| Headache                 | 0 (0)      | 6 (35)          |
| SAEs                     |            |                 |
| CAPS                     | 1 (5.9)    | 0 (0)           |
| Cryptorchidism           | 1 (5.9)    | 0 (0)           |
| Diarrhea                 | 1 (5.9)    | 0 (0)           |
| Vomiting                 | 1 (5.9)    | 1 (5.9)         |
| Influenza                | 1 (5.9)    | 0 (0)           |
| Lung infection           | 1 (5.9)    | 0 (0)           |
| Wound infection (staphylococcal) | 1 (5.9) | 0 (0)          |
| Femur fracture           | 1 (5.9)    | 0 (0)           |
| Conductive deafness      | 0 (0)      | 1 (5.9)         |
| Abdominal pain           | 0 (0)      | 1 (5.9)         |
| Papillitis               | 0 (0)      | 1 (5.9)         |
| Pneumonia                | 0 (0)      | 2 (11.8)        |
| Bronchitis               | 0 (0)      | 1 (5.9)         |
| Meningitis aseptic       | 0 (0)      | 1 (5.9)         |
| Limb injury              | 0 (0)      | 1 (5.9)         |
| Hematoma                 | 0 (0)      | 1 (5.9)         |

* Except where indicated otherwise, values are the number (%) of patients. AEs = adverse events; SAEs = serious AEs; CAPS = cryopyrin-associated periodic syndrome.
† Since study enrollment.
‡ Experienced by at least 6 patients (>35%).

in the core study 9 (53%) of the 17 patients had AEs that were suspected to be related to the study drug, with 2 cases each of bronchitis and molluscum contagiosum; all other drug-related AEs were reported as single cases. In the extension study, 16 patients experienced at least 1 AE, and 10 (59%) of the 17 patients had AEs that were suspected to be related to the study drug, with the most common (observed in at least 3 of the patients) being diarrhea, pneumonia, rhinitis, pyrexia, and cough.

In the core study, 4 (24%) of the 17 patients experienced at least 1 SAE; routine hospitalization for operative correction of congenital cryptorchidism (complicated by staphylococcal wound infection), diarrhea, vomiting, femur fracture, flare of CAPS disease activity, influenza, and lung infection were reported. None of the SAEs were reported for more than 1 patient. In the extension study, 8 (47%) of the patients experienced at least 1 SAE. Pneumonia was reported in 2 patients (12%), while all other SAEs were reported in 1 patient each. The proportion of patients with SAEs during the extension study is depicted in Table 3. There were no deaths or discontinuations due to AEs or SAEs during the entire study.

**DISCUSSION**

CAPS is a rare autoinflammatory disease, caused by mutations in the *NLRP3* gene, and is often challenging to diagnose due to unfamiliarity among many physicians, the rarity of the disease, overlapping symptoms with other diseases (27), and the presence of low-penetrance genetic variants and poly-morphisms (28). This study assessed the long-term efficacy and safety of canakinumab in pediatric patients with CAPS ≤5 years of age across 7 countries and demonstrated clinical efficacy over a 3-year period.

Overall, at least one complete response was achieved in the majority of the patients (9 of 10 patients <2 years of age and 7 of 7 patients >2 years of age) over the 56 weeks of the core study, and complete response was achieved in all patients by the end of the study. During the core study, no discontinuations were observed. In the extension study no patient discontinued due to AEs, but 2 patients discontinued due to an unsatisfactory therapeutic effect. These results are consistent with the findings of various studies demonstrating sustained efficacy of subcutaneous canakinumab administered every 8 weeks (21,29). Another phase III study (22) demonstrated clinically and serologically inactive disease in 30% of patients after 2 months and 60% of patients after 6 months of canakinumab treatment. In our study, 75% of the NOMID patients (3 of 4) and 42% of the MWS patients (5 of 12) required dose up-titration, emphasizing the need for higher doses in severely affected patients. In addition, the improvement in disease activity, as measured by physician global assessment of disease activity, that was observed during the core study was sustained until the end of the study. This improvement was also reflected in the assessment of skin disease (urticaria-like skin rash).

Although CRP levels decreased over time until the end of the study, such a pattern of change was not evident in SAA levels. This was due to the fact that although all patients had clinically active disease, several patients had low prestudy SAA levels. Additionally, underpowering and missing data probably contributed to this seemingly discrepant SAA response to treatment. In periodic fever syndromes such as CAPS, pretreatment SAA levels do not always capture systemic inflammation, even when patients have clinically active disease, as previously observed in other studies of children with CAPS (22).

No new safety signals were observed, and the safety profile was consistent with that reported in previous canakinumab studies (21). No deaths were reported during the study. There were few clinically significant findings on audiograms or neurologic or ophthalmologic assessments. In this study, all but 1 of the patients underwent MRI of the brain and inner ear, and all MRI findings were reported as normal. This is consistent with the results of
previous studies of the effect of canakinumab treatment on neurologic symptoms in CAPS patients (21,30). Although it has been suggested that anakinra may have superior efficacy for the amelioration of inflammatory cerebrospinal fluid (CSF) biomarkers in patients with NOMID based on observations in a small number of patients who underwent serial CSF examination while sequentially receiving anakinra or canakinumab (31), there are no hard data to suggest that this translates into worse neurologic outcomes for NOMID patients treated with canakinumab compared to those treated with anakinra. This is clearly an area that will require ongoing surveillance in the longer term. As compared with the β-Conﬁdent Registry (32) (which assessed 288 patients across 13 countries and different age groups, all treated with canakinumab), the frequency of AEs and SAEs was signiﬁcantly lower in the present study of very young pediatric patients; however, it is likely explained by small patient numbers in our study (and hence wider conﬁdence limits regarding this observation) compared with the β-Conﬁdent Registry data.

In our study, canakinumab did not show any negative impact on postvaccination antibody production following the administration of non-live vaccines in 9 pediatric patients. Recent studies have suggested that pneumococcal vaccines can trigger severe local and systemic inﬂammation (25) in CAPS patients, who are genetically prone to overactivation of inﬂammasomes (32). Hence, the potential beneﬁts of pneumococcal immunization need to be assessed against safety signals. Of the 3 patients who received pneumococcal vaccine in this study, none developed an inﬂammatory ﬂare. This study excluded inoculation with live attenuated vaccines because the current consensus is that these vaccines are contraindicated in patients treated with biologic agents. Interestingly, a 4-month-old patient with CAPS (NOMID) treated with canakinumab was recently reported to have been immunized with live attenuated vaccines (measles, rubella, varicella, and mumps) and achieved sufﬁcient antibody titer without any complications (33).

Given the rarity of CAPS and the vulnerable nature of this young pediatric patient population, this was a nonrandomized and nonblinded study of a pragmatically small sample size. Although an extension study was included, the follow-up period was still very limited for what is likely to be lifelong treatment. We acknowledge that the long-term impact of canakinumab treatment on hearing loss and other potential late sequelae remains uncertain. Ultimately, much longer phase IV registry type studies may provide more data on the impact of canakinumab over decades of treatment.

Overall, our results demonstrate that canakinumab is an effective treatment for patients with CAPS ≤5 years of age. Canakinumab appears to have no effect on the ability to produce antibodies against standard childhood non-live vaccines, and no vaccine-associated disease flares were observed. The safety proﬁle of canakinumab was thus acceptable and similar in this very young pediatric cohort to that observed in previous studies of older patients.

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AUTHOR CONTRIBUTIONS

All authors were involved in drafting the article or revising it critically for important intellectual content, and all authors approved the final version to be published. Dr. Brogan had full access to all of the data in the study and takes responsibility for the integrity of the data and the accuracy of the data analysis.

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ROLE OF THE STUDY SPONSOR

Novartis Pharma AG facilitated the study design, provided writing assistance for the manuscript, and reviewed and approved the manuscript prior to submission. The authors independently collected the data, interpreted the results, and had the final decision to submit the manuscript for publication. Writing assistance was provided by Deepak Pakalapati and Anupama Tantia (Novartis Healthcare Pvt. Ltd, India). Publication of this article was not contingent upon approval by Novartis Pharma AG.

ADDITIONAL DISCLOSURES

Author Wei is an employee of China Novartis Institutes for Biomedical Research Company, Ltd.

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