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Phase 3 Trials Promise Highly Effective Option for Hidradenitis Suppurativa

Milan – Two similarly designed phase 3 trials have associated a biologic targeted at interleukin-17A (IL-17A) with uncommon efficacy for control of hidradenitis suppurativa (HS). The trials, called SUNSHINE and SUNRISE, together comprise the largest set of phase 3 data in this disease ever assembled. Of those exposed to the IL-17A inhibitor, more than 40% of patients in both trials achieved the primary outcome of a meaningful reduction in the Hidradenitis Suppurativa Clinical Response (HiSCR) scale.

Based on the progress these data represent for HS, "I think looking back we will think of this as being an important tipping point toward progress in this incredibly debilitating disease," reported Dr. Alexa B. Kimball, Professor of Dermatology, Harvard Medical School, Boston, Massachusetts. Affiliated with the Department of Dermatology at Beth Israel Deaconess Medical Center, Dr. Kimball was the principal investigator of this phase 3 program and a co-author of the North American HS treatment guidelines published in 2019.

The study drug for the SUNSHINE and SUNRISE trials was secukinumab, an IL-17A inhibitor which is administered by subcutaneous injection. Both studies were identically designed and planned for pooled analysis. The data reported are from the first 16 weeks of follow-up of the 52-week trials. Enrollment was limited to patients with long-established moderate or severe disease. In this challenging population the

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challenging population, the primary endpoint was reached by about 45% of patients on active therapy versus slightly more than 30% of those on placebo.

Relative Benefit Achieved Within 2 Weeks

"There was a rapid early response seen at about 2 weeks with steeply increasing benefit to about 4 weeks and then continued benefit through the course of the 16-week follow-up," reported Dr. Kimball (Figure 1). As someone who specializes in the treatment of HS and

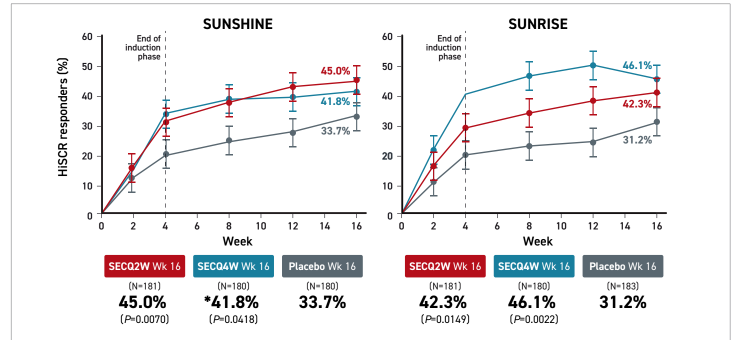


FIGURE 1 | SUNSHINE & SUNRISE Trials: HiSCR Achieved at Week 16

*Did not reach statistical significance.

One-sided nominal p-values are based on a logistic regression model, the primary estimand, and multiple imputation. Error bars represent 95% Cl.

CI, confidence intervals; HiSCR, hidradenitis suppurativa clinical response; N, number of patients in group; Q2W, every two weeks; Q4W, every four weeks; SEC, secukinumab 300 mg; Wk, week.

Adapted from Kimball AB et al. As presented during EADV 2022, D3T01.1A.

recognizes the difficulty of achieving a sustained level of control in a substantial proportion of patients, Dr. Kimball added, "I couldn't be more pleased to present these results."

The SUNSHINE trial enrolled 541 HS patients, and the SUNRISE trial enrolled 543. In each, patients were randomized in a 1:1:1 ratio to 300 mg of secukinumab every 2 weeks (Q2), 300 mg of secukinumab every 4 weeks (Q4), or placebo.

The primary outcome of a HiSCR clinical response was defined as >50% reduction in the number of inflammatory lesions with no increase in the number of abscesses or draining fistulae. In addition, patients with a baseline numerical rating scale (NRS) pain score of >3 (on a 0 to 10 scale) were required to achieve >30% reduction in pain from baseline.

Response Rates with Q2 and Q4 Doses Are Similar

All four of the active treatment arms in the two studies achieved a response rate of \geq 40%. For the 45% response rate in the Q2 arm (*P*=0.007) of SUNSHINE and the 46.1% response rate in the Q4 arm of SUNRISE (*P*=0.002), the differences relative to placebo were highly statistically significant. The response in the Q2 arm of SUNRISE also met statistical significance (*P*=0.0149). In the Q4 arm (*P*=0.418), the advantage for secukinumab relative to placebo fell just short of a predefined level of significance at 16 weeks, but Dr. Kimball pointed to imbalances in baseline severity that might have played a role. More will be understood as responses are tracked over the remaining 36 weeks of the study. Although the full 52-week data are now being analyzed and are not yet ready for presentation, Dr. Kimball provided a preview by saying, outcomes "are continuing to improve, and we are very excited."

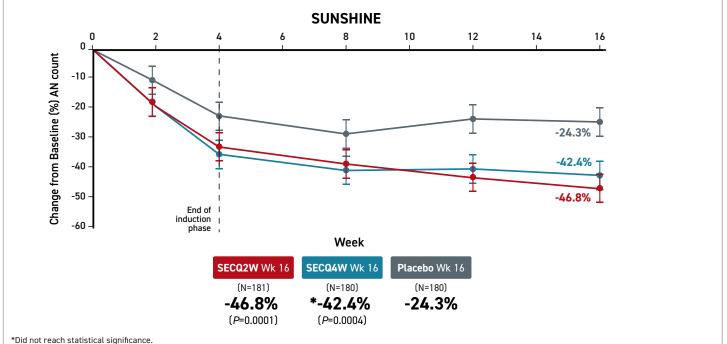
The efficacy of secukinumab against HS was reflected in all secondary endpoints, including substantial reductions in characteristic skin lesions and pain. These favourable effects are likely to explain the advantage of secukinumab for measures of quality of life at 16 weeks.

Flares, Nodules, and Abscesses All Reduced

"In both studies, the proportion of patients experiencing flares was lower with secukinumab compared to placebo at all timepoints from week 2 to week 16," Dr. Kimball reported. Similarly, the reduction in abscesses and inflammatory nodule count in both studies was apparent almost immediately, reaching significance by 2 weeks for both active treatment arms relative to placebo. The curves for these outcomes continued to separate over the course of follow-up with overall reductions about 2-fold greater in any secukinumab arm relative to placebo by 16 weeks (*P* values ranged from =0.0051 to <0.0001).

The same type of pattern was observed in curves tracking the rates of flares. A separation of the curves in both studies for active treatment arms relative to placebo was observed at 2 weeks and then increased over time. By 16 weeks, the Q2 dose relative to placebo was the most effective for suppressing flares (15.4% vs. 29.0% on placebo) in the SUNSHINE trial (Figure 2). While the Q4 dose was the most effective (15.6% vs. 27.0%) in the





One-sided nominal p-values are based on ANCOVA, the secondary estimand, and multiple imputation. Error bars represent SE.

AN, abscess and inflammatory nodule; ANCOVA, analysis of covariance; HS, hidradenitis suppurativa; N, number of patients in group; Q2W, every two weeks; Q4W, every four weeks;

SE, standard error; SEC, secukinumab 300 mg; Wk, week.

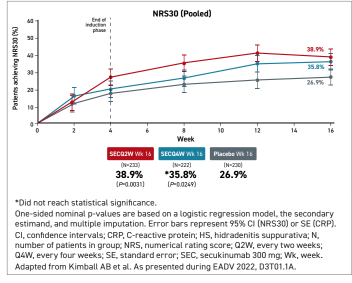
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SUNRISE trial, the other 2 doses also significantly lowered the risk of flares relative to placebo.

Pain Reduction Observed Within 2 Weeks

On control of pain, both the Q2 (P=0.0031) and the Q4 dose (P=0.0249) were significantly more effective when defined as >2 unit reduction in the Global Assessment of Skin Pain (GASP) validated tool. When graphed, the difference in the pooled analysis was apparent by week 2 with a further advantage achieved by the end of 16 weeks (Figure 3).





"I want to emphasize this benefit," Dr. Kimball said. "Pain has been a hard endpoint to meet in clinical trials, and we saw a reduction that was both statistically and clinically meaningful on secukinumab."

FIGURE 4 | Substantial Improvements Demonstrated in Quality-of-life Analyses

The reduction in disease burden was reflected in the quality of life analyses conducted with the Dermatology Life Quality Index (DLQI). For a clinically meaningful improvement in DLQI, defined as a >5 point reduction, scores were already markedly higher in both of the Pain has been a hard endpoint to meet in clinical trials, and we saw a reduction that was both statistically and clinically meaningful.

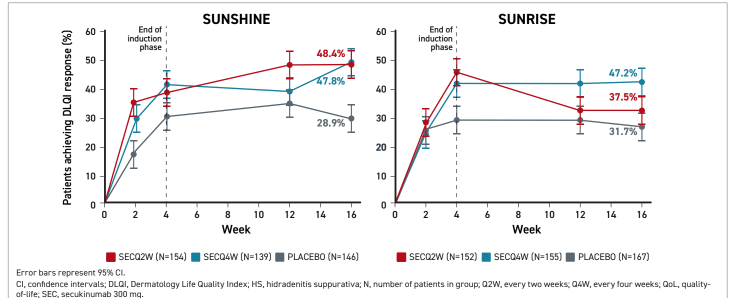
active treatment arms relative to placebo by 2 weeks in both trials. Although the further increase by week 4 was generally maintained in both arms in SUNRISE, quality of life scores climbed further by week 16 relative to week 4 in the SUNSHINE trial (Figure 4).

Representative Population Enrolled

The baseline demographics of the SUNSHINE and SUNRISE trials suggest that the population studied was representative. Although female representation (56.3%) was somewhat lower than that reported in other studies and surveys of HS, the mean age of 36.2 years is generally consistent with peak age of activity. About 60% of those enrolled had Hurley stage 2 at entry with the remaining presenting with Hurley stage 3. About 10% to 15% were on systemic antibiotics when enrolled. Prior experience with biologics was permitted and was reported by about 25% of patients. The mean body weight of those enrolled was more than 90 kg, reflecting the substantial representation by overweight and obese patients.

A history of HS for more than 1 year was an entry criterion, but, as is typical of the HS population, most reported that the onset of symptoms had taken place at least 7 years prior to the study despite the fact that the condition was not diagnosed in most until much more recently.

"A delay to the diagnosis is a common problem in HS, and the data from this trial in that regard is actually



Adapted from Kimball AB et al. As presented during EADV 2022, D3T01.1A.

encouraging," Dr. Kimball said. In studies she has conducted previously, diagnostic delay has generally been on the order of 10 years or longer, suggesting, "I think we are starting to do a little bit better in paying attention to this disorder."

HS Data Add to Study Drug Safety Record

Consistent with the experience in other disease treatments with secukinumab, which has indications for psoriasis and several forms of arthritis, the drug was well tolerated.

"There were no surprises in terms of safety. We know a lot about secukinumab from the previous studies in other diseases and the safety profile is very clean overall," Dr. Kimball said.

While adverse events overall and adverse events of special interest were generally consistent between those randomized to either dose of secukinumab or placebo, this was also reflected in discontinuation rates. The study retention rate was 93%, which Dr. Kimball suggested was "remarkable" when considering that the trial was conducted during the COVID-19 pandemic. The discontinuation rate for adverse events was less than 3% in all study arms.

While there was no difference overall in infections, "even the rate of Candida was not really different than placebo even in this population of relatively young women who are at high risk for vaginal candidiasis," Dr. Kimball reported.

Relative to many diseases, a 40-45% response rate for the primary endpoint of significant disease improvement might seem modest, but it is impressive in HS, according to Dr. Kimball. The most recent guidelines (Alikhan A. et al. *J Am Acad Dermatol* 2019;81:91-101) include a long list of potential therapies, which is needed because response rates to any given therapy are typically low. The heterogeneity of response to HS therapies is attributed to individual differences in contributing pathophysiology, which includes inflammatory processes and hormonal disturbances. Genetic susceptibility is likely to play a role in at least some patients, but it is now considered likely that there is no optimal therapy for all patients.

Biologics Are the Cornerstone of HS Therapy

"There is growing attention to phenotypes for directing treatment, particularly in early stages or when symptoms remain mild," Dr. Kimball said. However, she concurs with the published guidelines, which identify biologics "as the cornerstone of treatment for moderate-to-severe HS."

If these phase 3 trial data lead to secukinumab approval for HS, this IL-17A-targeted agent will join the tumour necrosis factor (TNF) inhibitor adalimumab as only the second approved biologic for this disease. It cannot be determined from the SUNSHINE and SUNRISE data if IL-17A is a better target than TNF to control HS-related inflammation, but these new data confirm that it is an effective target. More treatments for this disease are urgently needed, particularly those with moderate-to-severe manifestations.

"HS has become an incredibly active field right now in terms of understanding the underlying biology," said Dr. Kimball. By validating a new target for disease control, she considers the SUNSHINE and SUNRISE trial to be a "milestone" toward improved symptom control.

Conclusion

Two multicenter randomized phase 3 trials have validated IL-17A as an important therapeutic target in patients with HS. In two similarly designed trials, SUNSHINE and SUNRISE, the IL-17A inhibitor secukinumab, whether administered in an every-2- or every-4-week subcutaneous dose, significantly reduced skin involvement, reduced pain, and improved an overall measure of HS activity at the end of 16 weeks compared to placebo. The final results at 52 weeks are not yet available, but sustained benefit is expected. Even within 16 weeks, the effect of secukinumab, which was well tolerated, translated into improved quality of life.

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