International observational atopic dermatitis cohort to follow natural history and treatment course: TARGET-DERM AD study design and rationale

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BMJ Open  International observational atopic dermatitis cohort to follow natural history and treatment course: TARGET-DERM AD study design and rationale

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ABSTRACT

Introduction As new topical and systemic treatments become available for atopic dermatitis (AD), there is a need to understand how treatments are being used in routine clinical practice, their comparative effectiveness and their long-term safety in diverse clinical settings.

Methods and analysis The TARGET-DERM AD cohort is a longitudinal, observational study of patients with AD of all ages, designed to provide practical information on long-term effectiveness and safety unobtainable in traditional registration trials. Patients with physician-diagnosed AD receiving prescription treatment (topical or systemic) will be enrolled at academic and community clinical centres. Up to 3 years of retrospective medical records, 5 years of prospective medical records, and optional biological samples and patient-reported outcomes will be collected. The primary aims include characterisation of AD treatment regimens, evaluation of response to therapy, and description of adverse events.

Ethics and dissemination TARGET-DERM has been approved by a central IRB (Copernicus Group IRB, 5000 Centregreen Way Suite 200, Cary, North Carolina 27513) as well as local and institutional IRBs. No additional Ethics Committee reviews. Results will be reviewed by a publications committee and submitted to peer-reviewed journals.

Trial registration number NCT03661866, pre-results.

BACKGROUND

Atopic dermatitis is, by definition, a chronic condition that waxes and wanes over time. Defects of the epidermal barrier, immune dysregulation and environmental factors are thought to affect disease expression, and it is now recognised as a lifelong disposition with variable clinical manifestations. Despite being one of the leading contributors to non-fatal disease burden worldwide, very little is understood about long-term disease control.

Strengths and limitations of this study

- TARGET-DERM atopic dermatitis is a longitudinal cohort study of patients with atopic dermatitis, including up to 3 years of retrospective data and 5 years of prospective data, allowing for assessment of changes over time in individual patients.
- The study has broad inclusion criteria to allow for capture of patient populations that may be under-represented in clinical trials, and inclusion of patients from both academic and community practices will improve generalisability.
- The study relies on standardised extraction from routinely collected medical records.
- The study collects validated investigator global assessments and quarterly patient-reported outcome measures.
- The study has a biospecimen collection.

Pragmatic, real-world studies that capture long-term variability in atopic dermatitis disease activity and management are needed and can provide complementary data to clinical trials. Based on growing knowledge of immune pathways, the pharmacological options for management of atopic dermatitis have expanded over the past several years, with further expansion expected during the next decade. Clinical trials are often performed in highly selected, adherent study participants who lack significant comorbidities beyond the disease in question. These trials are a good measure of clinical efficacy, but the more germane question for clinicians and their patients is one of real-world effectiveness. Indeed, participants enrolled in dermatology clinical trials may not be highly representative of the general atopic dermatitis population. Subgroups that might be excluded or under-represented in clinical...
METHODS AND ANALYSIS

Study design

TARGET-DERM AD is a longitudinal, observational cohort study of atopic dermatitis. The primary aims of TARGET-DERM include characterisation of treatment regimens, evaluation of patient outcomes and description of adverse events. Secondary aims include evaluation of the relationship between atopic dermatitis and comorbid medical conditions. No specific treatments will be dictated by enrolment into the study; patient management will follow each site’s local standard of care.

Adult and paediatric patients of all ages will be enrolled from up to 100 practices, including both academic practices affiliated with a university health system and community-based or private practice clinical centres. The first patient was enrolled on 25 January 2019 and, as of March 2020, there are 34 active sites and an additional 10 in the start-up phase (Figure 1). Site recruitment is ongoing. Enrolment rates by site type, geography, patient age, patient race/ethnicity, socioeconomic status and type of treatment (topical vs systemic) are reviewed by the academic steering committee monthly and compared with national statistics. The recruitment strategy is designed to be adaptive to ensure adequate cohort diversity.

Study participants

As this study seeks to reflect real-world clinical practice, there are minimal inclusion/exclusion criteria. Patients of all ages being managed or treated for atopic dermatitis with at least one prescription treatment will be enrolled. A diagnosis of atopic dermatitis made by the treating physician, as in standard clinical practice, will be required for enrolment. During site orientation, diagnostic recommendations from the American Academy of Dermatology will be reviewed with enrolling physicians, and participants will be asked UK Working Party criteria at enrolment to enable post-hoc evaluations based on these diagnostic criteria. Patients will be excluded if they are unable to provide written informed consent or assent, or are participating in an investigational study of a systemic treatment for atopic dermatitis at the time of enrolment. Concurrent participation in investigational studies of topical treatments or other registries or observational studies on atopic dermatitis treatment outcomes, however, are permitted.

Patient and public involvement

At the time of consent, patients in TARGET-DERM are presented with joining this study as a way to advance research in their disease area. In addition, a non-profit medical specialty society dedicated to education and research on atopic dermatitis, the International Eczema Council, has joined the steering committee. Patients were not directly involved with the development or conduct of the study. The plan is to share study results through publication; posters and manuscripts will be displayed on the TARGET website.

Study procedures

After informed consent is obtained and eligibility confirmed, baseline demographic and background information will be collected. On enrolment and during each follow-up visit for atopic dermatitis, clinicians will record a validated investigator global assessment (vIGA-AD) score. Up to 3 years of retrospective medical records, and up to 5 years of prospective medical records will be collected biannually. Records will include clinic notes, hospitalisations, laboratory reports, telephone contact reports, medication lists, biopsy results and imaging. Structured variables to be extracted from these records include demographic factors, personal and family medical history, AD disease characteristics, current medications, laboratory tests and detailed AD treatment history (Box 1). There are no study-mandated interventions planned.

Participants will be asked to complete patient-reported outcome (PRO) assessments at baseline and every
3 months thereafter. These address itch, pain and sleep, quality of life, severity, work productivity and activity impairment, tailored to adult or paediatric populations (table 1). PROs were chosen based on Harmonizing Outcome Measures in Eczema clinical practice set recommendations, and core domains for participants on systemic and phototherapies will include those recommended by the international treatment of atopic dermatitis registry task force (TREAT) eDelphi exercise.

Optional biological samples will be collected yearly for future translational research on genetic and immunophenotypes and include blood, cutaneous tape strips and saliva. Participants will consent to each optional specimen. When possible, blood samples will be collected during regular clinical blood work.

The TARGET steering committee (currently composed of the academic coauthors on this manuscript) developed the study protocol and research plan and gave input on site selection and database design. The study is privately funded by TARGET PharmaSolutions and can be used to provide postauthorisation safety data as requested by licensing and regulatory bodies. TARGET stakeholders, including steering committee members, industry partners, regulatory representatives, and a nonprofit medical specialty society participate in steering committee meetings to drive the research agenda for TARGET studies. The steering committee has oversight over any public dissemination of the data generated in this study.

### Data management

Participating investigational sites provide medical records (including structured and unstructured data) from consented participants. All data will be processed and stored centrally via an electronic data capture system maintained by TARGET PharmaSolutions. Biosamples will be collected at the sites, sent for storage at a central biorepository and tracked in the electronic data capture system by TARGET; investigators and stakeholders can apply to use the samples for future research. Coding of data will be performed using standardised data extraction forms and MedDRA and WHODrug dictionaries. Data will be monitored for quality and missingness. Specific efforts will be made to investigate and document any reasons leading to premature study termination. Adverse events will be evaluated bi-annually and reported to regulatory bodies as outlined in the study safety monitoring plan.

### Study outcomes

The primary study outcomes are response to therapy and adverse events. Response to therapy will be based on changes in vIGA-AD as collected during standard care visits and change in patient-reported outcomes collected quarterly. Changes in therapy (ie, discontinuation, stepping-up, or stepping down) will also be considered. Adverse events will be defined as any untoward medical occurrence which deviates from, or is an exacerbation of, the subject’s medical history after the time of consent. Additional secondary outcomes include the occurrence and impact of comorbid medical conditions on treatment regimens and vice versa.

### Sample size

At least 4000 study participants from up to 100 clinical centres will be recruited for TARGET-DERM-AD. The sample size was chosen based on logistical considerations and the ability to address multiple study objectives; post-hoc calculations suggest that a sample size of 4000 AD participants (assuming 2000 per group) would enable detection of effect sizes as small as 0.07 for changes in the primary outcome of vIGA-AD.

### Discussion

TARGET-DERM AD is designed to enable the study of real-world treatment patterns and long-term outcomes in atopic dermatitis, including both comparative effectiveness and safety endpoints. The study has broad inclusion criteria to allow for capture of patient populations that may be underrepresented in clinical trials,
Table 1: Study measures and timing

<table>
<thead>
<tr>
<th>Activity</th>
<th>Prior 3 years*</th>
<th>Screening/entry visit</th>
<th>Follow-up Year 1</th>
<th>Follow-up Year 2</th>
<th>Follow-up Year 3</th>
<th>Follow-up Year 4</th>
<th>Follow-up Year 5</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>Month 0</td>
<td>Month 6</td>
<td>Month 12</td>
<td>Month 18</td>
<td>Month 24</td>
<td>Month 30</td>
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<td>Informed consent</td>
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<tr>
<td>Inclusion/exclusion</td>
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<td>Demographics and background forms</td>
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<td>Biologic sample collection</td>
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<tr>
<td>Patient reported outcome surveys†</td>
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<td></td>
<td></td>
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<td>Patient-reported outcomes will be completed every 3 months</td>
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<tr>
<td>Investigator global assessment†</td>
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<td></td>
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<td>To be collected at all standard of care visits</td>
</tr>
<tr>
<td>Medical records submission*</td>
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<td>X</td>
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</tbody>
</table>

For paediatrics, these include: UKWPC, POEM-Paediatric/Proxy, PROMIS-Itch Severity, NRS-Pain, NRS-Sleep, PROMIS-Paediatric Anxiety, PROMIS-Paediatric Depression, Children’s Dermatology Life Quality Index (CDLQI) and PO-SCORAD; these are all completed every 3 months except for UKWPC (one time) and POEM-Paediatric/Proxy, CDLQI and PO-SCORAD (every 6 months).

*Three years of historical records will be submitted following the screening/enrolment visit. During the follow-up period, medical records will be submitted every 6 months for up to 5 years. The first submission during the follow-up period will be 6 months following the screening/enrolment visit. Additional interim medical records submissions may be requested.

†UK Working Party diagnostic criteria (UKWPC), Patient-Oriented Eczema Measure (POEM), Patient-Reported Outcomes Measurement Information System (PROMIS)-Itch Severity, Numerical Rating Scale for Pain (NRS-Pain), NRS-Sleep, PROMIS-General, PROMIS-Mood and Sleep, PROMIS-Activity and Clothing, PROMIS-Scratching Behaviour, PROMIS-Anxiety, PROMIS-Depression, PROMIS-Itch Triggers, PROMIS-Itch Quality, Dermatology Life Quality Index (DLQI), Patient-Oriented-SCORing Atopic Dermatitis (PO-SCORAD) and the Work Productivity and Activity Impairment (WPAI). These are all completed every 3 months except for UKWPC, PROMIS-Itch Triggers and PROMIS-Itch Quality (one time), and DLQI and WPAI (every 6 months).

The protocol was approved by a central institutional review board (Copernicus IRB), and sites-specific IRB approvals are obtained prior to patient enrolment where required (see online supplemental file 1). Results will be published in peer-reviewed journals and presented at national and international meetings and reported in accordance with the Strengthening the Reporting of Observational Studies in Epidemiology recommendations.

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and recruitment of participants from a diverse group of academic and community practices will improve generalisability. TARGET-DERM AD fills an important niche; although other atopic dermatitis registries exist, these are largely focused on the impacts of specific treatments (Pediatric Eczema Elective Registry (PEER) and A Prospective Pediatric Longitudinal Evaluation to Assess the Long-Term Safety of Tacrolimus Ointment for the Treatment of Atopic Dermatitis (APPLE)), skin infections associated with AD (Atopic Dermatitis Research Network), or systemic treatments (TREAT). An important limitation is documentation of atopic dermatitis disease activity, which could be prone to inaccuracies when relying on real-world assessments. In particular, the frequency of patient visits and details in routinely collected medical records may not accurately capture intermittent disease activity and severity. Regular quarterly patient reported assessments and biosamples will enhance the internal validity for the subset of participants consenting to these additional outcomes.

TARGET-DERM AD will serve as the prototype for a larger cohort study of immune-mediated inflammatory skin conditions (IMISC), to ultimately include additional participants with psoriasis, hidradenitis suppurativa, vitiligo, and alopecia areata, and to enrol up to 15,000 participants. Although multiple registries and cohorts for psoriasis exist, such data sources are limited for other inflammatory skin diseases. Given the overlap in risk factors and treatment options for IMISC, a joint registry will benefit the growing understanding of these diseases, including the natural history, heterogeneity of different patient subpopulations, real-world management strategies and response to treatment. Additionally, ongoing monitoring of side effect profiles of emerging therapeutic agents used for multiple indications will enhance knowledge of the safety profile. Furthermore, the availability of an established cohesive research network allows nimble responses to investigations of new treatment paradigms with existing agents, as well as future generations of IMISC therapies.

ETHICS AND DISSEMINATION

The protocol was approved by a central institutional review board (Copernicus IRB), and sites-specific IRB approvals are obtained prior to patient enrolment where required (see online supplemental file 1). Results will be published in peer-reviewed journals and presented at national and international meetings and reported in accordance with the Strengthening the Reporting of Observational Studies in Epidemiology recommendations.
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Contributors KA, JMC and EG-Y: substantial contributions to conception or design of the work, drafting of the work or revising it critically, final approval of the version to be published and agreement to be accountable for all aspects of the work. JIS, ELS, ASP, LE, RB and DT: substantial contributions to conception or design of the work, drafting of the work or revising it critically, final approval of the version to be published. JK and JH: substantial contributions to conception or design of the work, final approval of the version to be published. LD and SEW: substantial contributions to the conception or design of the work, critical revisions, final approval of the version to be published.

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Competing interests KA: receives compensation for consulting services from TARGET PharmaSolutions; no other competing interests. JIS, AbbVie, AnaptyBiologics, Arena, Asana, Boehringer-Ingelheim, Dermira, DermaVant, DS Biopharma, Eli Lilly, Galderma, GliaxoSmithKline, Glenmark, Incyte, Kiniksa, LEO Pharma, Lunex, Menlo, Novartis, Pfizer, Regeneron Pharmaceuticals, Sanofi – consultant or advisory board member; Regeneron Pharmaceuticals, Sanofi – speaker; ELS: grants and personal fees from AbbVie, grants and personal fees from Eli Lilly, grants from Galderma, grants from Kyowa Hakko Kirin, grants from personal fees from LEO Pharmaceutical, grants from Merck, grants and personal fees from Pfizer, grants and personal fees from Regeneron, personal fees from Sanofi, personal fees from Dermira, grants from Galderma, grants and personal fees from MedImmune, grants from Novartis, grants from Tioga, grants from Celgene, personal fees from Boehringer-Ingelheim, personal fees from Dermavant, personal fees from Forto Bio, personal fees from Incyte, personal fees from Menlo Therapeutics, personal fees from Ortho Dermatologics, personal fees from Pierre Fabre Dermo Cosmetique, personal fees from Valeant. ASP: Investigator for AbbVie, AnaptyBiologics, Celgene, Eli Lilly, Galderma, Incyte, Leo, Janssen, Novartis, and Regeneron; consultant with honorarium for Amgen, Amgen, Asana, Boehringer-Ingelheim, Castle Creek, Celgene, Dermavant, Dermira, Eli Lilly, Exicure, Forte, Galderma, Lenus, Leo, MEDA Corp, Meiji Seika, Novan, Novartis, Pfizer, Regeneron, Sanofi-Genzyme, and Sol Gel. LE: Consultant/Speaker/Advisory Board: Almirall, DermaVant, Dermira, DS Biopharma, Forte, Galderma, Incyte, LEO, L’Oreal, Matsriys, Otsuka, Novartis, Ortho Dermatologics/Valeant, Pfizer/Anacor, Regeneron, Sanofi-Genzyme. Investigator: Abbvie, LEO, REGNERON, Sanofi-Genzyme. DSM: Asana, Ichnos, Glenmark. RB: Advisory Board Member, Consultant, Speaker, Investigator for and/ or receives honoraria or grant from AbbVie, AntibioTx, Arcutis, Arena Pharma, Asana Biosciences, Bellus Health, Boehringer-Ingelheim, DermaVant, Eli Lilly, EMD Serono, Galderma, Incyte, Kiniksa, Kyowa Kirin, Neekera, LEO Pharma, Novan, Pfizer, Relxlex, RAPT, Regeneron, Sanofi Genzyme and Sienna. Employee and shareholder of Innovarderm Research. JK: Personal fees from Novartis, Pfizer, Amgen, Lilly, Boehringer, BMS, Biogenidec, Janssen, AbbVie, Leo Pharma, ENSILUS, Menlo, Aristeo, Sanofi, Sun Pharma, Almiral, Arena, Ventyx, Aclaris, Galapagos. Grants paid to Institution from Novartis, Pfizer, Amgen, Lilly, Boehringer, Innovarderm, BMS, Janssen, AbbVie, Parexel, Leo Pharma, Vitae, Akros, Regeneron, Allergan, Novan, Biogen MA, Sienna, UCB, Celgene, Botanix, Incyte, Avidlim, Exicure. JH: consultant for Pfizer, Genzyme/Sanofi, Aclaris Therapeutics, Incyte, Theos Medicines, Sun Pharmaceuticals, LEO Pharma, Villaris Therapeutics, Dermavant, Tamprian, AbbVie, Inc., Janssen, TeVido BioDevices, EMD Serono, Almiral, Boston Pharma, Sonoma Biotherapeutics, Inc., Methuselah Health, Tbi Biotech, Pandion, Cogen Therapeutics, Inc., Admix and BridgeBio. Investigator for Pfizer, Genzyme/Sanofi, Aclaris Therapeutics, Incyte, Theos Medicines, Sun Pharmaceuticals, LEO Pharma, Villaris Therapeutics, Dermavant, AbbVie, Inc., TeVido BioDevices, EMD Serono and Pandion. Equity in TeVido Biodichos. Rough, Villaris Therapeutics, Inc. Scientific founder of Villaris Therapeutics, Inc. LD: employee at TARGET PharmaSolutions. SEW: employee at TARGET PharmaSolutions. JMC: employee at TARGET PharmaSolutions. DT: is a lecturer and/or consultant for AbbVie, Almirall, Amgen, Asana Biosciences, Biogen Idec, BiocAD, Boehringer Ingelheim, Bristol- Myers Squibb, Celgene, DS-Biopharma, GliaxoSmithKline, Glenmark, Kyowa Kirin, Leo Pharma, Eli Lilly, Novartis, Regeneron, Sandoz, Sanofi-Aventis and UCB, and received grants from AbbVie and Novartis (paid to institution). EGY: employee of Mount Sinai and has received research funds (grants paid to the institution) from AbbVie, Almirall, Amgen, AnaptyBio, Asana Biosciences, Boehringer-Ingelheim, Celgene, DermaVant, DS Biopharma, Eli Lilly, Galderma, Ichnos Sciences, Innovarderm, Janssen, Kiniksa, Kyowa Kirin, Leo Pharma, Novan, Pfizer, Realexar, Regeneron, Sienna Biopharma, UCB, and Union Therapeutics. EGY is also a consultant for AbbVie, Almirall, Amgen, Arena Pharmaceuticals, Asana Biosciences, AstaZeneca Biopharmaceuticals, Boehringer-Ingelheim, Cara Therapeutics, Celgene, Concert, DBV, Derma, DS Biopharma, Eli Lilly, EMD Serono, Escalier, Galderma, Ichnos Sciences, Kyowa Kirin, Leo Pharma, Mitsubishi Tanabe, Pandion Therapeutics, Pfizer, RAPT Therapeutics, Regeneron, Sanofi, Sienna Biopharma, and Union Therapeutics.

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Harmonising outcome measures for eczema. Available: www.homeforeczema.org


