A PHASE 3, OPEN-LABEL, MULTICENTER STUDY OF ALXN1210 IN CHILDREN AND ADOLESCENTS WITH ATYPICAL HEMOLYTIC UREMIC SYNDROME (aHUS)

Unique Protocol ID: ALXN1210-aHUS-312
NCT Number: NCT03131219
EudraCT Number: 2016-002499-29
Date of SAP: 07 November 2018
A PHASE 3, OPEN-LABEL, MULTICENTER STUDY OF ALXN1210 IN CHILDREN AND ADOLESCENTS WITH ATYPICAL HEMOLYTIC UREMIC SYNDROME (aHUS)
1. APPROVAL SIGNATURES

Date dd Mmm yyyy

Date dd Mmm yyyy

Date dd Mmm yyyy
# TABLE OF CONTENTS

1. APPROVAL SIGNATURES .......................................................................................2  
2. TABLE OF CONTENTS, LIST OF TABLES, AND LIST OF FIGURES .................3  
3. LIST OF ABBREVIATIONS AND DEFINITIONS OF TERMS .........................8  
4. DESCRIPTION OF THE PROTOCOL ...................................................................9  
   4.1. Changes from Analyses Specified in the Protocol ........................................11  
   4.2. Changes from Analyses Specified in the Previous Version of the SAP .........11  
   4.3. Planned Analyses .........................................................................................12  
5. DEFINITIONS ......................................................................................................13  
   5.1. Efficacy ........................................................................................................13  
   5.1.1. Primary Endpoint(s) .................................................................................13  
   5.1.2. Secondary Endpoints ...............................................................................13  
   5.1.3. Exploratory Endpoints .............................................................................14  
   5.1.4. Other Efficacy Endpoints .........................................................................14  
   5.2. Safety ..........................................................................................................14  
   5.2.1. Adverse Events .........................................................................................14  
   5.2.2. Vital Signs ...............................................................................................15  
   5.2.3. Laboratory Assessments ...........................................................................15  
   5.2.4. Physical Examinations .............................................................................16  
   5.2.5. Electrocardiograms ...............................................................................16  
   5.2.6. Immunogenicity .......................................................................................16  
   5.2.7. Other Safety Events of Special Interest ..................................................16  
6. DATA SETS ANALYZED (STUDY POPULATIONS) ........................................17  
   6.1. Full Analysis Set ..........................................................................................17  
   6.2. Per Protocol Set ..........................................................................................17  
   6.3. Safety Set ....................................................................................................18  
   6.4. Other Sets ....................................................................................................18  
7. STATISTICAL ANALYSIS ...............................................................................19  
   7.1. Study Patients ..............................................................................................19  
   7.1.1. Disposition of Patients ............................................................................19
| 7.1.2 | Protocol Deviations | 19 |
| 7.1.3 | Demographics, Disease Characteristics, and History | 20 |
| 7.1.3.1 | Demographics | 20 |
| 7.1.3.2 | Disease Characteristics | 20 |
| 7.1.3.3 | Medical History and Baseline Physical Examination | 20 |
| 7.1.4 | Prior and Concomitant Medications / Nonpharmacologic Procedures | 20 |
| 7.2 | Efficacy Analyses | 21 |
| 7.2.1 | Primary Analysis | 21 |
| 7.2.1.1 | Handling of Dropouts or Missing Data | 22 |
| 7.2.1.2 | Subgroup Analysis | 22 |
| 7.2.1.3 | Multicenter Studies | 22 |
| 7.2.1.4 | Hypothesis Testing and Significance Level | 22 |
| 7.2.1.5 | Sensitivity Analyses | 23 |
| 7.2.2 | Secondary Analyses | 23 |
| 7.2.2.1 | Dialysis Requirement Status | 23 |
| 7.2.2.2 | Time to Complete TMA Response | 23 |
| 7.2.2.3 | Complete TMA Response Status Over Time and Alternative Definitions | 23 |
| 7.2.2.4 | eGFR Value and Change from Baseline | 24 |
| 7.2.2.5 | CKD Stage | 24 |
| 7.2.2.6 | Hematologic Parameters | 25 |
| 7.2.2.7 | Hemoglobin Response | 25 |
| 7.2.2.8 | Quality of Life | 25 |
| 7.2.3 | Exploratory Analyses | 26 |
| 7.2.4 | Other Efficacy Analyses | 26 |
| 7.2.5 | Pharmacokinetic and Pharmacodynamic Analyses | 26 |
| 7.3 | Safety Analyses | 26 |
| 7.3.1 | Study Duration, Treatment Compliance, and Exposure | 26 |
| 7.3.2 | Adverse Events (AEs) | 27 |
| 7.3.2.1 | Overall Summary of Adverse Events | 27 |
| 7.3.2.2 | AEs and SAEs by System Organ Class (SOC) and Preferred Term (PT) | 27 |
| 7.3.2.3 | AEs and SAEs by SOC, PT, and Relationship | 27 |
| 7.3.2.4 | AEs and SAEs by SOC, PT, and Severity | 28 |
| 7.3.2.5 | Deaths, Other SAEs, and Other Significant Adverse Events | 28 |
7.3.3. Other Safety ................................................................................................................28
7.3.3.1. Analyses for Laboratory Tests ....................................................................................28
7.3.3.2. Vital Signs ..................................................................................................................28
7.3.3.3. Other Safety Parameters of Special Interest ...............................................................28
8. REFERENCES ...........................................................................................................30
9. APPENDICES ............................................................................................................31
9.1. Protocol Schedule of Events .......................................................................................31
9.2. Changes from Analyses Specified in the Previous Version of the SAP .....................31
9.3. Sample Size, Power, and Randomization ....................................................................31
9.4. Technical Specifications for Derived Variables ...................................................................31
9.4.1. Age ..............................................................................................................................31
9.4.2. Definition of Baseline Values .....................................................................................31
9.4.3. Change from Baseline .................................................................................................31
9.4.4. Analysis Visits ............................................................................................................31
9.4.5. Analysis Value ............................................................................................................32
9.4.6. Adverse Events ...........................................................................................................32
9.4.7. Adverse Events Related to Meningococcal Infection .................................................33
9.4.8. Concomitant Medications/Therapies ........................................................................33
9.4.9. Serum Creatinine and eGFR .......................................................................................33
9.4.10. Platelets and Hemoglobin ...........................................................................................33
9.4.11. Assignment of CKD Stage ..........................................................................................34
9.5. Questionnaires and Patient Reported Outcomes .........................................................35
9.5.1. Pediatric FACIT-Fatigue ............................................................................................35
9.5.2. Resource Utilization Patient Questionnaire ...............................................................36
9.5.3. Patient-Reported AHUS Symptoms Questionnaire ..................................................37

LIST OF TABLES

Table 1: Abbreviations and Acronyms ..................................................................................8
Table 2: Adverse Event Severity Grading Scale .....................................................................14
Table 3:  Causality Assessment Descriptions .................................................................15
Table 4:  Adverse Events (AEs) and Serious AEs ......................................................27
Table 5:  Age and Reference Date ............................................................................31
Table 6:  Stages of Chronic Kidney Disease .............................................................34
LIST OF FIGURES

Figure 1: Study Design Schematic for Clinical Protocol ALXN1210-aHUS-312 .................. 9
3. LIST OF ABBREVIATIONS AND DEFINITIONS OF TERMS

The following abbreviations and acronyms are used in this Statistical Analysis plan (SAP).

Table 1: Abbreviations and Acronyms

<table>
<thead>
<tr>
<th>Abbreviation or acronym</th>
<th>Explanation</th>
</tr>
</thead>
<tbody>
<tr>
<td>ADA</td>
<td>Antidrug antibodies</td>
</tr>
<tr>
<td>ADAMTS13</td>
<td>A disintegrin and metalloproteinase with a thrombospondin type 1 motif, member 13</td>
</tr>
<tr>
<td>ADL</td>
<td>Activities of daily living</td>
</tr>
<tr>
<td>AEs</td>
<td>Adverse events</td>
</tr>
<tr>
<td>aHUS</td>
<td>Atypical hemolytic uremic syndrome</td>
</tr>
<tr>
<td>BP</td>
<td>Blood pressure</td>
</tr>
<tr>
<td>C5</td>
<td>Complement component 5</td>
</tr>
<tr>
<td>CDF</td>
<td>Cumulative distribution function</td>
</tr>
<tr>
<td>CI</td>
<td>Confidence interval</td>
</tr>
<tr>
<td>CKD</td>
<td>Chronic kidney disease</td>
</tr>
<tr>
<td>CSR</td>
<td>Clinical Study Report</td>
</tr>
<tr>
<td>CTCAS</td>
<td>Common Terminology Criteria for Adverse Events</td>
</tr>
<tr>
<td>ECG</td>
<td>Electrocardiogram</td>
</tr>
<tr>
<td>eCRF</td>
<td>Electronic Case Report Form</td>
</tr>
<tr>
<td>eGFR</td>
<td>Estimated Glomerular Filtration Rate</td>
</tr>
<tr>
<td>EQ-5D-3L</td>
<td>EuroQol 5 dimensions 3 level</td>
</tr>
<tr>
<td>FACT</td>
<td>Functional assessment of chronic illness therapy</td>
</tr>
<tr>
<td>FAS</td>
<td>Full Analysis Set</td>
</tr>
<tr>
<td>IV</td>
<td>Intravenous</td>
</tr>
<tr>
<td>LDH</td>
<td>Lactate dehydrogenase</td>
</tr>
<tr>
<td>LLT</td>
<td>Lowest level term</td>
</tr>
<tr>
<td>LOCF</td>
<td>Last Observation Carried Forward</td>
</tr>
<tr>
<td>MedDRA</td>
<td>Medical Dictionary for Regulatory Activities</td>
</tr>
<tr>
<td>mITT</td>
<td>Modified intent to treat</td>
</tr>
<tr>
<td>MMRM</td>
<td>Mixed model for repeated measures</td>
</tr>
<tr>
<td>PD</td>
<td>Pharmacodynamic</td>
</tr>
<tr>
<td>PE/PI</td>
<td>Plasma exchange/plasma infusion</td>
</tr>
<tr>
<td>PK</td>
<td>Pharmacokinetic</td>
</tr>
<tr>
<td>PT</td>
<td>Preferred Term (MedDRA)</td>
</tr>
<tr>
<td>PTAEs</td>
<td>PreTreatment Adverse Events</td>
</tr>
<tr>
<td>QoL</td>
<td>Quality of life</td>
</tr>
<tr>
<td>q8w</td>
<td>Once every 8 weeks</td>
</tr>
<tr>
<td>SAEs</td>
<td>Serious adverse events</td>
</tr>
<tr>
<td>SAS®</td>
<td>Statistical Analysis Software®</td>
</tr>
<tr>
<td>SAP</td>
<td>Statistical Analysis Plan</td>
</tr>
<tr>
<td>SD</td>
<td>Standard deviation</td>
</tr>
<tr>
<td>SOC</td>
<td>System Organ Class (MedDRA)</td>
</tr>
<tr>
<td>SOP</td>
<td>Standard operating procedure</td>
</tr>
<tr>
<td>TEAEs</td>
<td>Treatment-Emergent Adverse Events</td>
</tr>
<tr>
<td>TMA</td>
<td>Thrombotic microangiopathy</td>
</tr>
</tbody>
</table>
4. DESCRIPTION OF THE PROTOCOL

Study ALXN1210-aHUS-312 is a Phase 3, open-label, single-treatment arm, multicenter study to evaluate the safety efficacy, PK, and PD of ALXN1210 administered by intravenous (IV) infusion in approximately 23 to 28 pediatric patients, from birth to < 18 years of age, with confirmed diagnosis of aHUS. The study has 2 cohorts. Cohort 1 includes complement inhibitor treatment-naïve patients; Cohort 2 includes eculizumab-experienced adolescent patients (12 to < 18 years of age). The study consists of 3 periods: a Screening Period (of up to 7 days for Cohort 1 or up to 28 days for Cohort 2), a 26-week Initial Evaluation Period, and an Extension Period of up to 2 years.

Figure 1 illustrates the study design.

Figure 1: Study Design Schematic for Clinical Protocol ALXN1210-aHUS-312

Abbreviations: q4w = once every 4 weeks; q8w = once every 8 weeks

* The Screening Period is up to 7 days for complement inhibitor treatment-naïve patients (ie, Cohort 1) and up to 28 days for eculizumab-experienced adolescent patients (ie, Cohort 2).

Approximately 23 to 28 pediatric (< 18 years of age) patients with documented aHUS are planned for this study. The minimum number of patients for each age category is as follows:

- Birth to < 2 years: 4 patients
- 2 to < 6 years: 4 patients
- 6 to < 12 years: 4 patients
- 12 to < 18 years: 8 patients

Cohort 2 patients must be 12 to < 18 years of age.
The primary objective of the study is to assess the efficacy of ALXN1210 in complement inhibitor treatment naïve pediatric patients (ie, Cohort 1) with aHUS to inhibit complement-mediated thrombotic microangiopathy (TMA) as characterized by thrombocytopenia, hemolysis, and renal impairment.

Secondary objectives for complement inhibitor treatment-naïve patients (ie, Cohort 1) include the following:

1. To characterize the safety and tolerability of ALXN1210
2. To evaluate the efficacy of ALXN1210 by the following additional measures:
   a. Dialysis requirement status
   b. Time to Complete TMA Response
   c. Complete TMA Response status over time
   d. Observed value and change from baseline in estimated glomerular filtration rate (eGFR)
   e. Chronic kidney disease (CKD) stage, as evaluated by eGFR at select target days and classified as improved, stable (no change), or worsened compared to baseline
   f. Observed value and change from baseline in hematologic parameters (platelets, lactate dehydrogenase (LDH), hemoglobin)
   g. Increase in hemoglobin of $\geq 20$ g/L from baseline, observed at two separate assessments obtained at least 4 weeks (28 days) apart, and any measurement in between
   h. Change from baseline in quality of life (QoL), as measured by Pediatric Functional Assessment of Chronic Therapy (FACIT) Fatigue questionnaires
3. To characterize the pharmacokinetic (PK)/pharmacodynamics (PD) of ALXN1210:
   a. Changes in serum ALXN1210 concentration over time
   b. Changes in serum free complement component 5 (C5) concentrations over time
4. To evaluate the long term safety and efficacy of ALXN1210

The secondary objectives for eculizumab-experienced patients (ie, Cohort 2) are to assess the following:

1. To characterize the safety and tolerability of ALXN1210
2. To evaluate the efficacy of ALXN1210 by the following measures:
   a. Dialysis requirement status
   b. Observed value and change from baseline in eGFR
   c. CKD stage, as evaluated by eGFR at select target days and classified as improved, stable (no change), or worsened compared to baseline
   d. Observed value and change from baseline in hematologic parameters (platelets, LDH, hemoglobin)
   e. Change from baseline in QoL, as measured by Pediatric FACIT Fatigue questionnaire
3. To characterize the PK/PD of ALXN1210:
   a. Changes in serum ALXN1210 concentration over time
b. Changes in serum free C5 concentrations over time

4. To evaluate the long-term safety and efficacy of ALXN1210

For additional details, refer to the protocol.

4.1. Changes from Analyses Specified in the Protocol

None

4.2. Changes from Analyses Specified in the Previous Version of the SAP

The original SAP (dated 27 March 2017) was amended once. A summary of the changes is presented below.

- In Section 7.1.2, removed the specification for production of a table presenting the number and percentage of patients meeting or not meeting inclusion/exclusion criteria. A listing is deemed sufficient.
- In Section 5.2.3, added text specifying that local laboratory results may be used for analysis when no central laboratory results are available. Text was also added to this section to address in vitro erythrocyte lysis referred to as table top hemolysis (TTH) caused by sample mishandling.
- In Section 7.2.2.3 and Section 7.2.2.7, an example indicating patients to be included in a summary of response by visit was deleted as it was incorrectly implying that certain patients would be excluded from some time points.
- In Section 7.2.1.2 and Section 7.3, age groupings for subgroups analyses of efficacy and safety data were adjusted.
- In Section 9.4.10, added text indicating that the exclusion from analysis of hemoglobin and platelets results when concurrent with blood transfusion is only applicable to postbaseline assessments. Because pre-treatment assessments, even if concurrent with blood transfusion, need to meet the inclusion criteria.
- In Section 4, the description of the protocol was adjusted to primarily reflect the addition of eculizumab-experienced adolescent patients to the study. This adjustment resulted in the description of 2 separate cohorts; complement inhibitor treatment-naïve patients (ie, Cohort 1), and eculizumab-experienced adolescent patients (ie, Cohort 2). Section 5, Section 6 and Section 7 were adjusted to specify definitions and analyses applicable to each cohort.
- In Section 4.3, text was adjusted to include an additional interim analysis.
- In Section 7.2.5, a description of basic summaries of pharmacokinetic and pharmacodynamic data was added.
- In Section 5.1.1, added text specifying that local laboratory results may be used for analysis when no central laboratory results are available.
4.3. Planned Analyses

An interim analysis is planned when 12 to 14 complement inhibitor treatment-naïve (ie, Cohort 1) patients have completed or withdrawn from the 26-week Initial Evaluation Period. A second interim analysis is planned after all study patients have completed or withdrawn from the 26-week Initial Evaluation Period.

Additionally, a final analysis to summarize long-term efficacy, safety, and PK parameters will be performed at the end of the 2-year Extension Period.

The present document serves to describe these planned analyses.
5. DEFINITIONS

5.1. Efficacy

All efficacy endpoints are applicable to both cohorts unless otherwise noted.

5.1.1. Primary Endpoint(s)

The primary efficacy endpoint of the study (only applicable to Cohort 1) is Complete TMA Response during the 26 week Initial Evaluation Period. The criteria for Complete TMA Response are the following:

1. Normalization of platelet count (defined as platelet count ≥ 150,000 per microliter)
2. Normalization of LDH (defined as LDH ≤ upper limit of normal)
3. ≥ 25% improvement in serum creatinine from baseline

Patients must meet all Complete TMA Response criteria at 2 separate assessments obtained at least 4 weeks (28 days) apart, and any measurement in between.

If local laboratory tests are used for LDH, platelet count, and serum creatinine, duplicate samples will be collected for central laboratory testing to ensure baseline and postbaseline measurements for analyses are resulted from the central laboratory. In the event of duplicate samples from local and central laboratories (for any time point), central laboratory results will be used for analysis. Local laboratory results will only be used for analysis if no central laboratory results are available.

5.1.2. Secondary Endpoints

The secondary efficacy endpoints of the study are the following:

5. Dialysis requirement status
6. Time to Complete TMA Response (only applicable to Cohort 1)
7. Complete TMA Response status over time (only applicable to Cohort 1)
8. Observed value and change from baseline in eGFR
9. CKD stage, as evaluated by eGFR at select target days and classified as improved, stable (no change), or worsened compared to baseline
10. Observed value and change from baseline in hematologic parameters (platelets, LDH, hemoglobin)
11. Increase in hemoglobin of ≥ 20 g/L from baseline, observed at 2 separate assessments obtained at least 4 weeks (28 days) apart, and any measurement in between (only applicable to Cohort 1)
12. Change from baseline in QoL, as measured by Pediatric FACIT Fatigue questionnaires
5.1.3. Exploratory Endpoints

Patients will report signs or symptoms of aHUS and resource utilization using the Patient-reported aHUS Symptoms Questionnaire and the Resource Utilization Patient Questionnaire, respectively. These are only applicable to Cohort 1 patients enrolled under a protocol version preceding amendment 5, as they will no longer be collected under protocol amendment 5.

The Investigator will evaluate extra-renal signs or symptoms of aHUS using a composite of clinical laboratory measurements, vital signs, and an organ system review. This is applicable to both Cohort 1 and Cohort 2.

5.1.4. Other Efficacy Endpoints

Not applicable

5.2. Safety

The safety and tolerability of ALXN1210 will be evaluated by physical examinations, vital signs, electrocardiograms (ECGs), laboratory assessments, and incidence of adverse events (AEs) and serious adverse events (SAEs). The proportion of patients who develop antidrug antibodies (ADA) will also be assessed. Safety endpoints are applicable to both Cohort 1 and Cohort 2.

5.2.1. Adverse Events

Treatment-emergent AEs (TEAEs) are defined as any event not present prior to exposure to Investigational Product or any event already present that worsens in either intensity or frequency following exposure to Investigational Product. An AE will be considered a TEAE if the start date and time is on or after the start date and time of the first study drug infusion. All other AEs are considered PreTreatment Adverse Events (PTAEs).

The severity of AEs will be graded using Common Terminology Criteria for Adverse Events (CTCAE) version 4.03 or higher. A grading (severity) scale is provided for each AE term. Each LLT will be coded to a MedDRA preferred term.

Grade refers to the severity of the AE. The CTCAE assigns a grade of 1 through 5, with unique clinical descriptions of severity for each AE (Table 2).

Table 2: Adverse Event Severity Grading Scale

<table>
<thead>
<tr>
<th>Grade</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Grade 1</td>
<td>Mild; asymptomatic or mild symptoms; clinical or diagnostic observations only; intervention not indicated</td>
</tr>
<tr>
<td>Grade 2</td>
<td>Moderate; minimal, local or noninvasive intervention indicated; limiting age-appropriate instrumental activities of daily living (ADL)&lt;sup&gt;a&lt;/sup&gt;</td>
</tr>
<tr>
<td>Grade 3</td>
<td>Severe or medically significant, but not immediately life-threatening; hospitalization or prolongation of hospitalization indicated; disabling; limiting self-care ADL&lt;sup&gt;b&lt;/sup&gt;</td>
</tr>
<tr>
<td>Grade 4</td>
<td>Life-threatening consequences; urgent intervention indicated.</td>
</tr>
<tr>
<td>Grade 5</td>
<td>Death related to AE.</td>
</tr>
</tbody>
</table>

Abbreviations: ADL = activities of daily living; AE = adverse event

<sup>a</sup> Instrumental ADL refers to preparing meals, shopping for groceries or clothes, using the telephone, managing money, etc.

<sup>b</sup> Self-care ADL refers to bathing, dressing and undressing, feeding self, using the toilet, taking medications, and not bedridden.
An Investigator must provide a causality assessment (Unrelated, Unlikely, Possible, Probable, or Definite) for all AEs (both serious and nonserious) based upon the Investigator’s medical judgment and the observed symptoms associated with the event (Table 3).

<table>
<thead>
<tr>
<th>Assessment</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Not Related/Unrelated</td>
<td>Suggests that there is no causal association between the investigational product and the reported event.</td>
</tr>
<tr>
<td>Unlikely Related</td>
<td>Suggests that the clinical picture is highly consistent with a cause other than the investigational product but attribution cannot be made with absolute certainty and a relationship between the investigational product and AE cannot be excluded with complete confidence.</td>
</tr>
<tr>
<td>Possibly Related</td>
<td>Suggests that treatment with the investigational product may have caused or contributed to the AE (ie, the event follows a reasonable temporal sequence from the time of drug administration and/or follows a known response pattern to the investigational product, but could also have been produced by other factors).</td>
</tr>
<tr>
<td>Probably Related</td>
<td>Suggests that a reasonable temporal sequence of the event with the investigational product administration exists and the likely causal association of the event with the investigational product. This will be based upon the known pharmacological action of the investigational product, known or previously reported adverse reactions to the investigational product or class of drugs, or judgment based on the Investigator’s clinical experience.</td>
</tr>
<tr>
<td>Definitely Related</td>
<td>Temporal relationship to the investigational product, other conditions (concurrent illness, concurrent medication reaction, or progression/expressio of disease state) do not appear to explain event, corresponds with the known pharmaceutical profile, improvement on discontinuation, reappearance on rechallenge.</td>
</tr>
</tbody>
</table>

For more details see Section 11.7 of the protocol, and Section 9.4 of this SAP.

### 5.2.2. Vital Signs

Vital sign measurements will be taken after the patient has been resting for at least 5 minutes, and will include systolic and diastolic blood pressure (BP) (millimeters of mercury [mmHg]), pulse oximetry, heart rate (beats/minute), respiratory rate (breaths/minute), and oral or tympanic temperature (degrees Celsius [°C] or degrees Fahrenheit [°F]). Physical growth (height, weight and head circumference) will be assessed.

### 5.2.3. Laboratory Assessments

Laboratory assessments are defined in section 11.4 of the protocol. Laboratory values will be graded according to the National Cancer Institute CTCAE version 4.03 or higher. Parameters which are not present in the CTCAE document will not be graded.

Samples collected at Screening may be tested at either a local or central laboratory. If local laboratory tests are used for LDH, platelet count, hemoglobin, and serum creatinine, duplicate samples will be collected for central laboratory testing to ensure baseline and postbaseline measurements for analyses are resulted from the central laboratory. In the event of duplicate samples from local and central laboratories (for any time point), central laboratory results will be
used for analysis. Local laboratory results will only be used for analysis if no central laboratory results are available.

It has been observed in other studies that up to 1% of central laboratory chemistry samples undergo in vitro erythrocyte lysis referred to as table top hemolysis (TTH) caused by sample mishandling. Due to the artefactual increase in LDH in samples affected by TTH, the potassium, ALT, AST, magnesium, phosphorous, and LDH values in samples affected by TTH will not be used in the analysis of any efficacy or safety endpoints. TTH samples from the central lab will be defined as having serum potassium $\geq 6$ mmol/L and LDH $\geq 2x$ ULN, and will be excluded from analyses as described above.

5.2.4. Physical Examinations

A physical examination will include the following assessments: general appearance; skin; head, ear, eye, nose, and throat; neck; lymph nodes; chest; heart; abdominal cavity; limb; central nervous system; and musculoskeletal system. An abbreviated physical examination consists of a body system relevant examination based upon Investigator judgment and patient symptoms.

5.2.5. Electrocardiograms

For each patient, single 12-lead digital ECGs will be collected according to the Schedule of Assessments. Patients must be supine for approximately 5 to 10 minutes before ECG collection and remain supine but awake during ECG collection.

The Investigator or designee will be responsible for reviewing the ECG to assess whether the ECG is within normal limits and to determine the clinical significance of the results. These assessments will be indicated on the CRF. For any clinically significant abnormal ECG results, the Investigator must contact the Medical Monitor to discuss the patient’s continued eligibility to participate in this protocol.

5.2.6. Immunogenicity

Blood samples will be collected to test for presence and titer of ADAs to ALXN1210 in serum prior to study drug administration as indicated in the Schedule of Assessments.

5.2.7. Other Safety Events of Special Interest

Meningococcal infections will be considered as AEs of special interest.
6. DATA SETS ANALYZED (STUDY POPULATIONS)

6.1. Full Analysis Set

The full analysis set (FAS) will be used for the primary analysis of efficacy.

The FAS for Cohort 1 will be based on a modified intent to treat (mITT) approach. With this approach, confirmation of eligibility in patients may occur after receiving study drug. Based on the above, the FAS will include all patients who receive at least 1 dose of ALXN1210, have at least 1 postbaseline efficacy assessment, and meet all of the following criteria:

- Patients who satisfy Inclusion Criterion number 2c
- Patients who satisfy Exclusion Criterion number 1
- Patients who satisfy Exclusion Criterion number 2

See protocol sections 8.1 and 8.2 for the full list and description of the inclusion and exclusion criteria.

The FAS for Cohort 2 will include all patients who receive at least 1 dose of ALXN1210 and have at least 1 postbaseline efficacy assessment.

6.2. Per Protocol Set

The per protocol set (PPS) for Cohort 1 will consist of FAS patients who meet all of the following criteria:

- Received 100% of the planned number of infusions during the 26 week Initial Evaluation Period
- Did not take any prohibited medications or undergo any prohibited procedures (see protocol section 9.7)
- Satisfied inclusion criteria 2 and 8 (or 10 for protocol amendment 5), and exclusion criteria 3, 10, 11, 12, 13, 15, 16, 17, 18, 21, 22 and 26 (or 28 for protocol amendment 5)

The per protocol set (PPS) for Cohort 2 will consist of FAS patients who meet all of the following criteria:

- Received 100% of the planned number of infusions during the 26 week Initial Evaluation Period
- Did not take any prohibited medications or undergo any prohibited procedures (see protocol section 9.7)
- Satisfied protocol amendment 5 inclusion criteria 3, 4 and 10, and exclusion criteria 3, 10, 11, 12, 15, 16, 17, 18, 21, 23, 24 and 28
6.3. Safety Set

Safety analyses will be performed on the Safety Set defined as all patients who receive at least 1 dose of study drug for both Cohort 1 and Cohort 2.

6.4. Other Sets

Not applicable.
7. **STATISTICAL ANALYSIS**

All analyses will be performed using Statistical Analysis Software® (SAS®) release, version 9.4 or higher (SAS Institute Inc., Cary, NC, USA) or other validated statistical software. Continuous variables will be summarized using descriptive statistics, including number of observations, mean, standard deviation (SD), median, minimum, and maximum values. Categorical variables will be summarized by frequency counts and percentage.

The analyses for complement inhibitor treatment-naïve patients (ie, Cohort 1) and eculizumab-experienced adolescent patients (ie, Cohort 2) will be conducted and reported separately. Analyses specific to Cohort 1 are indicated as such below; all other endpoints will be performed for both cohorts. Tabulated summaries will not include a direct comparison between Cohort 1 and Cohort 2.

7.1. **Study Patients**

7.1.1. **Disposition of Patients**

All patients will be included in the summaries of patient disposition, which will describe the frequency and percentage of patients enrolled and treated and who completed or withdrew from the study, along with reason for withdrawal from the study. The number and percentage of patients who are treated, discontinue treatment (along with reason for treatment discontinuation), complete or withdraw from the Initial Evaluation Period (along with reason for withdrawal), enter the Extension Period, and complete or withdraw from the Extension Period (along with reason for withdrawal) will be tabulated.

These summaries will be presented for both Cohort 1 and Cohort 2.

7.1.2. **Protocol Deviations**

Important protocol deviations will be determined per the standard operating procedure (SOP) “Identification, Handling, and Documentation of Protocol Deviations” (SOP-G-CL-0044). The number and percent of patients with specific protocol deviations will be summarized for all enrolled patients by major and minor deviations.

To ensure completeness of the list of important protocol deviations, the following will be identified programmatically from the database and shared with the Clinical Operations team:

1. Patients from whom informed consent was not obtained
2. Patients who violated any inclusion/exclusion criteria
3. Patients who took prohibited medications or underwent any prohibited procedure
4. Patients who received less than 100% of the protocol specified number of doses of study drug during the Initial Evaluation Period
5. Patients found not to satisfy one or more of the following inclusion/exclusion criteria (Inclusion Criterion number 2 or Exclusion Criteria numbers 1 and 2) whom are not discontinued from the study.
A by-patient listing of inclusion/exclusion criteria, as well as protocol deviations will be presented, separately.

This summary will be presented for both Cohort 1 and Cohort 2.

7.1.3. **Demographics, Disease Characteristics, and History**

Patient demographic and baseline characteristics, including medical history and transplant history, will be summarized for the FAS and Safety Set for both Cohort 1 and Cohort 2.

7.1.3.1. **Demographics**

The following demographic variables will be summarized:

- Gender
- Race
- Age (years) at first Infusion
- Weight at first Infusion
- Height at first Infusion

A by-patient listing of demographics will be generated.

7.1.3.2. **Disease Characteristics**

The following disease characteristics will be summarized:

- Age at first aHUS symptoms
- Pretreatment extra-renal signs or symptoms of aHUS
- A disintegrin and metalloproteinase with a thrombospondin type 1 motif, member 13 (ADAMTS-13) activity

A by-patient listing of disease characteristics will be generated.

7.1.3.3. **Medical History and Baseline Physical Examination**

Medical history will be summarized by number and percentage of patients within each Body System. Patients will be counted only once at the Body System level.

A separate table summarizing history of emergency room visits and hospitalizations due to aHUS, a table summarizing kidney transplant history, as well as a table presenting baseline physical examination results will also be generated.

By-patient listings of these data will be generated.

7.1.4. **Prior and Concomitant Medications / Nonpharmacologic Procedures**

Prior and concomitant medications (including vitamins and herbal preparations) will be coded using the World Health Organization Drug Dictionary (WHO-DRUG), while prior and
concomitant nonpharmacologic therapies and procedures (any therapeutic intervention, such as surgery/biopsy or physical therapy) will be coded using MedDRA.

Prior medications or procedures are defined as any non-study medications or procedures that were started, and were stopped, prior to the date of first infusion. Concomitant medications or procedures are defined as any non-study medications or procedures that were taken or occurred while the patient also received study medication. Prior and concomitant summaries will be presented separately.

Medications will be summarized by Anatomic-Therapeutic-Chemical (ATC) level 4 class and preferred drug name using frequency counts and percentages of patients for the Safety Set. Procedures will be summarized similarly, but by MedDRA Class and preferred term.

A table summarizing pretreatment plasma exchanges and plasma infusion as well as kidney dialysis will be generated. A similar table will summarize concomitant kidney dialysis.

In the event a patient receives a prohibited medication or undergoes a prohibited procedure, this will be reported as a protocol deviation as described in Section 7.1.2.

By-patient listings of these data will be provided separately.

These summaries will be presented for both Cohort 1 and Cohort 2.

7.2. Efficacy Analyses

The FAS is the primary analysis set for all efficacy analyses. The primary analysis and selected secondary efficacy analyses will be repeated on the PPS as sensitivity analyses.

7.2.1. Primary Analysis

The primary efficacy endpoint is Complete TMA Response during the 26-week Initial Evaluation Period. The primary analysis will consist in estimating the proportion of complete TMA responders among ALXN1210 treated patients. This will be performed by calculating the point estimate and a 2-sided 95% confidence interval (CI) for the proportion of complete TMA responders in ALXN1210 treated patients, where the numerator will be the number of patients achieving complete TMA response during the 26-week Initial Evaluation Period and the denominator will be the number of patients in the FAS. The CI will be based on exact confidence limits using the Clopper-Pearson method.

The criteria for Complete TMA response are presented in Section 5.1.1. Patients must meet all Complete TMA Response criteria at 2 separate assessments obtained at least 4 weeks (28 days) apart, and any measurement in between. To be considered a responder during the 26-week Initial Evaluation Period, the latest time point a patient can first meet the response criteria is 28 days before the Day 183 assessment.

Formal statistical comparison analyses are not planned for this estimation study. Results from ALXN1210 treated patients will be evaluated in the context of results observed in patients treated with eculizumab in prospective eculizumab studies.

This analysis will only be performed for Cohort 1.
7.2.1.1. **Handling of Dropouts or Missing Data**

For evaluation of Complete TMA Response during the 26-week Initial Evaluation Period (primary endpoint), patients missing an efficacy assessment that is part of the definition of Complete TMA Response while still on-study, will have their last observation carried forward (LOCF). For patients who will have discontinued from the study prior to Week 26, their data up to the time of discontinuation will be used to assess Complete TMA Response. A confirmatory result cannot be from an assessment that was carried forward from the initial assessment when all Complete TMA Response criteria were met.

For laboratory data, in the event of duplicate samples from local and central laboratories (for any time point), central laboratory results will be used for analysis.

7.2.1.2. **Subgroup Analysis**

The primary efficacy analysis will be repeated separately by the following subgroups within Cohort 1:

- gender (male, female)
- race
- ethnicity
- geographic region (by country)
- age at enrollment categories (birth to < 2 years, 2 to <6 years, 6 to <12 years, 12 to <18 years)
- kidney transplant history (yes, no)
- immunogenicity status (ever positive, always negative)
- dialysis within 5 days prior to treatment initiation (yes, no)

Given that the number of patients in these subgroups may be limited, the CIs will be based on exact confidence limits using the Clopper-Pearson method. For the same reason, some subgroup categories may be combined.

7.2.1.3. **Multicenter Studies**

The number of patients at each center is expected to be very small. Therefore, there will be no summarization of efficacy analyses by site.

7.2.1.4. **Hypothesis Testing and Significance Level**

This is an estimation study and no formal statistical tests are planned. The primary efficacy endpoint will be analyzed using an exact95% 2-sided confidence interval for the proportion of complete TMA responders.
7.2.1.5. Sensitivity Analyses

The primary and selected secondary efficacy endpoints will be analyzed based on the on the PPS to determine if the results are materially different from the analysis based on the FAS.

7.2.2. Secondary Analyses

Secondary Analyses will be performed on the FAS. Separate by-patient listings will be created for all secondary efficacy analysis parameters.

Continuous variables will be summarized using descriptive statistics, including number of observations and mean, standard deviation, median, minimum, and maximum values. Categorical variables will be summarized by frequency counts and percentage of patients.

7.2.2.1. Dialysis Requirement Status

An analysis will present the number and proportion of patients that are requiring dialysis over time. A 2-sided 95% CI for the proportion will be provided. The CI will be based on exact confidence limits using the Clopper-Pearson method. The summary will be produced for patients requiring dialysis within 5 days prior to ALXN1210 treatment initiation, not requiring dialysis within 5 days prior to ALXN1210 treatment initiation, and overall. A patient will be considered as not requiring dialysis at a specific postbaseline time point if they have been dialysis free for at least 5 days prior to that time point. A by-patient figure showing dialysis status or events over time will be presented. These summaries will be presented for both Cohort 1 and Cohort 2.

7.2.2.2. Time to Complete TMA Response

For the secondary efficacy endpoint of time to Complete TMA Response, Kaplan Meier cumulative distribution curves will be generated along with 2-sided 95% CIs. The corresponding summary table will present the cumulative distribution function (CDF) estimate, the number of patients at risk, the number of patients responding, and the number of patients censored at each postbaseline time point. The table will also present first quartile, median, and third quartile, along with corresponding 2-sided 95% CI, of time to complete response.

Complete TMA response is defined in Section 5.1.1. Patients must meet all Complete TMA Response criteria at two separate assessments obtained at least 4 weeks (28 days) apart, and any measurement in between. The time of the event of a confirmed complete TMA response will be considered as the first time point at which all the criteria for complete TMA response were met. Patients that did not have a response will be censored at the date of last visit or study discontinuation at the time when the analysis is performed. This analysis will only be performed for Cohort 1.

7.2.2.3. Complete TMA Response Status Over Time and Alternative Definitions

Complete TMA Response will also be summarized over time by presenting the number and proportion of responders along with a 2-sided 95% CI for each postbaseline time point. This summary will be accompanied by a similar presentation for the components of complete TMA response: proportion of patients with platelet count normalization, proportion of patients with LDH normalization, proportion of patients with hematologic normalization (platelet count and LDH), proportion of patients with ≥ 25% improvement in serum creatinine from baseline. The CI
will be based on exact confidence limits using the Clopper-Pearson method. These tabular summaries will be accompanied by a bar chart graphical figure displaying for each time point a bar for Complete TMA Response and each component in its definition, where the bar will represent the proportion of patients meeting that specific criterion.

A patient will be included in the analysis for a specific postbaseline time point if it was possible for the result at that time point to be confirmed or to serve as a confirmatory result.

A set of summaries similar to the ones described above will be produced without any requirement for a confirmatory result. This summary will be strictly based on a patient meeting the criterion at a specific time point.

A slightly modified version of complete TMA response will also be evaluated. The modification will only affect patients considered to be on dialysis at baseline (ie. patients requiring dialysis within 5 days prior to ALXN1210 treatment initiation). For these patients, the criterion requiring an improvement from baseline of 25% or more in serum creatinine, will be replaced by a postbaseline change in dialysis status (from requiring dialysis at baseline to no longer requiring dialysis) that is maintained for at least 4 weeks. A set of summaries similar to the ones described above will be produced based on this alternative definition.

The tabular summary of complete TMA response over time will also be performed on the PPS.

This analysis will only be performed for Cohort 1.

7.2.2.4. eGFR Value and Change from Baseline

Kidney function evaluated by eGFR will be summarized at baseline and each postbaseline time point using descriptive statistics for continuous variables for the observed value as well as the change from baseline. A value for eGFR will be imputed for patients requiring dialysis for acute kidney injury (see Section 9.4.9 for details). This summary will be repeated by the kidney transplant status at enrollment, and on the PPS.

A mixed model for repeated measures (MMRM) will be used to improve the precision of estimation of changes in eGFR over time. The model will include the fixed, categorical effect of visit and fixed, continuous effect of the baseline value as covariates. An unstructured covariance structure will be used to model the within patient errors. If this analysis fails to converge, the following structures will be tested and the final covariance structure will be determined by selecting the one with the lowest Akaike’s information criterion (AIC): first-order autoregressive, compound symmetry, and Toeplitz. The Kenward-Roger approximation will be used to estimate denominator degrees of freedom. The estimation of change from baseline on defined weeks alongside with 95% confidence intervals will be provided.

Figures presenting the observed values and the model based values over time will be provided.

This analysis will be performed for both Cohort 1 and Cohort 2.

7.2.2.5. CKD Stage

CKD stage will be summarized over time by presenting the number and proportion of patients that improved (excluding those with Stage 1 at baseline as they cannot improve), worsened (excluding those with Stage 5 at baseline as they cannot worsen), and stayed the same compared to CKD stage at baseline. Stage 5 will be considered the worst category, while stage 1 will be
considered the best category (see Section 9.4.11 for details). A 2-sided 95% CI for each proportion will be provided. The CIs will be based on exact confidence limits using the Clopper-Pearson method. This summary will be repeated by the kidney transplant status at enrollment. This analysis will be performed for both Cohort 1 and Cohort 2.

7.2.2.6. **Hematologic Parameters**

Hematologic parameters (platelets, LDH, hemoglobin) will be summarized at baseline and each postbaseline time point using descriptive statistics for continuous variables for the observed value as well as the change from baseline. This summary will be repeated on the PPS. As suggested in Section 7.2.2.4, a mixed model for repeated measures (MMRM) will be used to improve the precision of estimation of changes over time.

Figures presenting the observed values and the model based values over time for these hematologic parameters will be provided.

This analysis will be performed for both Cohort 1 and Cohort 2.

7.2.2.7. **Hemoglobin Response**

The number and proportion of patients with an increase from baseline in hemoglobin $\geq 20$g/L, observed at 2 separate assessments obtained at least 4 weeks (28 days) apart, and any measurement in between will be summarized over time by presenting the number and proportion of responders along with a 2-sided 95% CI for each postbaseline time point.

A patient will be included in the analysis for a specific postbaseline time point if it was possible for the result at that time point to be confirmed (ie, the patient has to be in the study for a subsequent confirmatory assessment at least 28 days later).

A set of summaries similar to the ones described above will be created without any requirement for a confirmatory result. This summary will be strictly based on a patient meeting the criterion at a specific time point.

This analysis will only be performed for Cohort 1.

7.2.2.8. **Quality of Life**

Quality of life will be evaluated in patients $\geq 5$ years of age by the using Pediatric FACIT Fatigue Questionnaire (patient-reported for patients who were $\geq 8$ years of age at the time of enrollment; caregiver-reported or caregiver assistance for patients who were $< 8$ years of age at the time of enrollment). This will be summarized at baseline and each postbaseline time point using descriptive statistics for continuous variables for the observed value as well as the change from baseline. As suggested in Section 7.2.2.4, a mixed model for repeated measures (MMRM) will be used to improve the precision of estimation of changes over time.

The number and proportion of patients with a 3 point improvement from baseline in the Pediatric FACIT Fatigue score will be summarized over time by presenting the number and proportion of patients along with a 2-sided 95% CI for each postbaseline time point.

A figure presenting the observed values and the model based values over time will be provided.

This analysis will be performed for both Cohort 1 and Cohort 2.
7.2.3. **Exploratory Analyses**

Resource Utilization Questionnaire (all patients) data will be summarized at each postbaseline assessment using discrete data summary statistics.

Patient-reported aHUS Symptoms Questionnaire (all patients) data will be summarized at each postbaseline assessment by presenting the number and proportion of patients with a specific symptom present. Extra-Renal Signs and Symptoms of aHUS data will be presented similarly.

These data will also be presented in detailed by-patient listings.

The Extra-Renal Signs and Symptoms of aHUS analysis will be performed for both Cohort 1 and Cohort 2, while the Resource Utilization Questionnaire and the Patient-reported aHUS Symptoms Questionnaire analysis will only be performed for Cohort 1.

7.2.4. **Other Efficacy Analyses**

Not applicable.

7.2.5. **Pharmacokinetic and Pharmacodynamic Analyses**

Individual serum concentration data for all patients from the FAS and who have evaluable PK data will be used to derive PK parameters for ALXN1210.

Graphs of mean serum ALXN1210 concentration-time profiles will be constructed. Graphs of serum concentration-time profiles for individual patients may also be provided. Actual dose administration and sampling times will be used for all calculations. Descriptive statistics will be calculated for serum concentration data at each sampling time, as appropriate.

Pharmacodynamic analyses will be performed for all patients from the FAS and who have evaluable PD data. The PD effects of ALXN1210 will be evaluated by assessing the absolute values and changes and percentage changes from baseline in serum free C5 serum concentrations over time, as appropriate. Descriptive statistics will be calculated for the PD data at each sampling time, as appropriate.

These analyses will be performed for both Cohort 1 and Cohort 2.

7.3. **Safety Analyses**

All safety analyses will be conducted on the Safety Set. All safety data will be provided in patient listings. AEs will be coded in MedDRA version 18 or higher and presented by MedDRA system organ class (SOC) and preferred term. All safety analyses will be presented by age category (0-5, 6-17 years of age) except for immunogenicity and physical examination summaries. No formal hypothesis testing is planned.

7.3.1. **Study Duration, Treatment Compliance, and Exposure**

Two summary tables of the extent of exposure to ALXN1210 will be presented. The duration on treatment (in weeks), the number of infusions, and the total dose (in mg) through the first 26 weeks (ie, Initial Evaluation Period) will be summarized using descriptive statistics. In addition, a similar table will be produced for exposure throughout the entire study.
7.3.2. Adverse Events (AEs)

The following definitions will be used for AEs:

- Pretreatment adverse events: Any AE that starts after providing informed consent, but before the first infusion of study drug.
- Treatment-emergent adverse event: Any AE that starts during or after the first infusion of study drug.
- Treatment-emergent SAE: A treatment-emergent AE (TEAE) that is serious.

Pretreatment AEs will be summarized in listings and TEAEs will be tabulated and presented as described below. These analyses will be performed for both Cohort 1 and Cohort 2.

7.3.2.1. Overall Summary of Adverse Events

An overall summary of AEs and SAEs will be presented. The number of events (n), and number and percentage of patients with events (n, %) will be shown for the following event subcategories:

Table 4: Adverse Events (AEs) and Serious AEs

<table>
<thead>
<tr>
<th>Relationship</th>
<th>Severity: Toxicity</th>
</tr>
</thead>
<tbody>
<tr>
<td>Related AEs (Possibly, Probably, or Definitely Related)</td>
<td>Grade 1</td>
</tr>
<tr>
<td>Not Related AEs (Not Related or Unlikely Related)</td>
<td>Grade 2</td>
</tr>
<tr>
<td></td>
<td>Grade 3</td>
</tr>
<tr>
<td></td>
<td>Grade 4</td>
</tr>
<tr>
<td></td>
<td>Grade 5</td>
</tr>
</tbody>
</table>

Fatal TEAEs, TEAEs resulting in study treatment discontinuation, TEAEs resulting in withdrawal from the study, and TEAEs that start during study drug administration will also be included in this overall summary. These statistics will be prepared separately for all AEs and SAEs.

Detailed listings of all AEs, SAEs, and related AEs, will be presented. These listings will include severity and relationship to treatment, as well as action taken regarding study treatment, other action taken, and AE outcome.

7.3.2.2. AEs and SAEs by System Organ Class (SOC) and Preferred Term (PT)

The number of TEAEs, and the number and percentage of patients with events will be presented by SOC and PT. Patients will be counted once in each SOC and PT. Percentages will be based on the total number of treated patients (safety set). SOCs will be listed in descending order of frequency of occurrence. SAEs will be summarized similarly.

7.3.2.3. AEs and SAEs by SOC, PT, and Relationship

The number of TEAEs, and the number and percentage of patients with events will be presented by SOC and PT as described above by relationship (related, not related). If a patient has more
than one occurrence of an AE, the strongest relationship to study treatment will be used in the relationship to treatment summary table. SAEs will be summarized similarly.

7.3.2.4. **AEs and SAEs by SOC, PT, and Severity**

The number of TEAEs, and the number and percentage of patients with events will be presented by SOC and PT as described above by severity (using toxicity grade). If a patient has more than one occurrence of an AE, the most severe occurrence of the AE will be used in the severity summary table. SAEs will be summarized similarly.

7.3.2.5. **Deaths, Other SAEs, and Other Significant Adverse Events**

Individual listings will be presented for AEs leading to study treatment discontinuation, AEs leading to withdrawal from the study, AEs starting during study drug administration, and fatal AEs.

In addition, a summary table and a listing of AEs related to meningococcal infections will be provided. See Section 9.4.7 for a list of AE preferred terms that will be considered for these summaries of meningococcal infections.

7.3.3. **Other Safety**

7.3.3.1. **Analyses for Laboratory Tests**

Observed values and changes from baseline in clinical chemistry, hematology, and urinalysis results will be summarized descriptively at baseline, and at each postbaseline time point. For laboratory results that can be classified as normal, low, or high based on normal range values, shifts from baseline in classification will be summarized for all study visits.

All data will be presented in listings, and a specific listing of abnormal results will be provided. For analysis purposes, laboratory results based upon standardized units will be used.

This analysis will be performed for both Cohort 1 and Cohort 2.

7.3.3.2. **Vital Signs**

Observed values as well as changes from baseline in body weight, height, head circumference and vital signs (blood pressure, heart rate, respiratory rate, pulse oxymetry, and temperature) at each time assessment will be summarized descriptively. A listing of vital signs data will be presented by patient, vital sign, and visit. This analysis will be performed for both Cohort 1 and Cohort 2.

7.3.3.3. **Other Safety Parameters of Special Interest**

7.3.3.3.1. **Electrocardiograms**

By-patient data listings of ECG parameters will be provided. Electrocardiograms will be evaluated and summarized as normal, abnormal not clinically significant, or abnormal clinically significant. A shift from baseline to worst on-study ECG table will be presented for ECG results. Observed values and change from baseline in ECG intervals (PR, RR, QT, and QTc) will be summarized descriptively at baseline and each postbaseline time point. QT interval will be
corrected for heart rate using Fridericia’s formula (QTcF). This analysis will be performed for both Cohort 1 and Cohort 2.

7.3.3.3.2. Immunogenicity

The number and percentage of patients with positive titers for ADAs to ALXN1210 and different titer categories will be summarized over time. The proportion of patients ever positive and the proportion of patients always negative will also be summarized. This analysis will be performed for both Cohort 1 and Cohort 2.

7.3.3.3.3. Physical Exam

Physical Exam parameters collected include General Appearance, Skin, HEENT (head, ears, eyes, nose and throat), Neck, Lymph Nodes, Chest, Heart, Abdominal Cavity, Limbs, Musculoskeletal, and Central Nervous System. Each parameter is assessed by the Investigator as Normal, Abnormal, or Not Examined. A listing of Physical Exam results will be presented by patient, Physical Exam parameter, and visit. This analysis will be performed for both Cohort 1 and Cohort 2.
8. REFERENCES


9. APPENDICES

9.1. Protocol Schedule of Events
See protocol section 7.3.

9.2. Changes from Analyses Specified in the Previous Version of the SAP
Not applicable.

9.3. Sample Size, Power, and Randomization
At least 16 pediatric patients (and no more than 24) will be enrolled in this study. This sample size is deemed appropriate to get proper representation in each of the 4 planned age groups (birth to < 2 years, 2 to < 6 years, 6 to < 12 years, 12 to < 18 years) and provide adequate safety information and precision level for the planned estimation.

9.4. Technical Specifications for Derived Variables

9.4.1. Age
Table 5: Age and Reference Date

<table>
<thead>
<tr>
<th>AGE</th>
<th>REFERENCE DATE</th>
</tr>
</thead>
<tbody>
<tr>
<td>• Age at Enrollment</td>
<td>• Date of Informed Consent Form Signing</td>
</tr>
<tr>
<td>• Age at First Symptoms</td>
<td>• Date of First Symptoms</td>
</tr>
<tr>
<td>• Age at First Infusion</td>
<td>• Date of First Infusion</td>
</tr>
</tbody>
</table>

9.4.2. Definition of Baseline Values
For the analysis, the baseline value is defined as the average of the values from the assessments performed prior to the first study drug infusion (these can include results from Screening and the Day 1 visit).

When a patient is on dialysis at baseline, then the first valid creatinine value to be used as the baseline value will be the first assessment ≥ 7 days post-dialysis. If a patient is on dialysis during the entire 26 week Initial Evaluation Period, then the baseline creatinine will not be calculated.

9.4.3. Change from Baseline
Change from baseline will be calculated as the baseline value subtracted from the value at a particular time point. If one of the values is missing and there are no pre-specified missing value imputation rules (see Section 7.2.1.1), then a change from baseline will not be calculated.

9.4.4. Analysis Visits
Summaries over postbaseline time points or analyses at specific postbaseline time points will be performed based on the list of visits described in the schedule of assessment of the protocol. For all assessments, the number of days from baseline will be calculated using the following
formula: (date of assessment) - (date of first study treatment) + 1. This number of days will be used to assign analysis visit. This may not always correspond to the electronic case report form (eCRF) visit.

The analysis visit assignment for a specific assessment will be based on visit windows around each scheduled visit for that specific assessment. The windows for each scheduled visit will go from the midpoint (in days) between the current visit and the previous scheduled visit to the midpoint between the current visit and the subsequent scheduled visit. If the interval separating 2 scheduled visits is an even number of days, that middle day will be included in the later visit window and excluded from the prior visit window. For example, for an assessment with a scheduled visit Day 127, and a prior scheduled visit Day of 113 and subsequent scheduled visit Day of 141, the window will start at 120 days from baseline and will go to 133 days from baseline.

9.4.5. Analysis Value

The values being considered for analysis at a specific postbaseline time point will be based on the analysis visit assigned to that value. If there is more than one non-missing value for a specific assessment with the same analysis visit, the value used for analysis will be the one for which the calculated number of days from baseline is closest to the scheduled visit day. If two values have the same analysis visit and are the same distance away from the scheduled visit day, the earlier of the 2 values will be used for analysis.

9.4.6. Adverse Events

The analysis of Adverse Events is described in detail in Section 7.3.2. Treatment-emergent AEs (TEAEs) are events with start dates and start times on or after the date and time of the first study drug dose. If the start date of an AE is partially or completely missing and the end (stop) date and time of the AE does not indicate that it occurred prior to first dose, then the determination of treatment-emergent status will be based on the following:

- If the start year is after the year of the first study drug dose, then the AE is treatment-emergent; else,
- If the start year is the same as the year of the first study drug dose and
  - the start month is missing, then the AE is treatment emergent; else if
  - the start month is present and is the same or after the month of the first study drug dose, then the AE is treatment-emergent; else,
- If the start date is completely missing, then the AE is treatment-emergent.

All other AEs are considered Pretreatment Adverse Events (PTAEs).

Related AEs are defined as possible, probable or definitely related. Unrelated AEs are defined as unlikely or not related.
9.4.7. **Adverse Events Related to Meningococcal Infection**

To find meningococcal events, the adverse event dataset will be searched for the following MedDRA Preferred Terms: Meningitis meningococcal, Meningococcal bacteraemia, Meningococcal infection, Meningococcal sepsis, Meningococcal carditis, Encephalitis meningococcal, Endocarditis meningococcal, Myocarditis meningococcal, Optic neuritis meningococcal, and Pericarditis meningococcal. In addition, a medical review will be done to ensure that no relevant events were missed.

9.4.8. **Concomitant Medications/Therapies**

The analysis of concomitant medications and therapies is described in detail in Section 7.1.4.

Concomitant medications or therapies are defined as any non-study medications or therapies that were taken or given while the patient also received study medication. A medication or therapy will be considered concomitant if the start date is on or after the date of the first study drug infusion, or if the start date is before the first infusion date and the end (stop) date is after the first infusion date. If the start date of a medication/therapy is partially or completely missing and the end (stop) date of the medication/therapy does not indicate that it ended prior to first infusion, then the determination of the concomitant status will be based on the following:

- If the start year is after the year of the first study drug infusion, then the medication/therapy is concomitant; else,
- If the start year is the same as the year of the first study drug infusion and
  - the start month is missing, then the medication/therapy is concomitant; else if
  - the start month is present and is the same or after the month of the first study drug infusion, then the medication/therapy is concomitant; else,
- If the start date is completely missing, then the medication/therapy is concomitant.

All other medications/therapies are considered Prior Medications/Therapies.

9.4.9. **Serum Creatinine and eGFR**

Serum creatinine measurements are not reliable with concurrent dialysis. Therefore, all serum creatinine values obtained while a patient is on dialysis will be excluded from all analyses. Estimated Glomerular Filtration Rate (eGFR) will be imputed with a value of 10 (in mL/min/1.73 m²) while a patient is on dialysis. A patient will be considered on dialysis from the first day of dialysis through 7 days after the end of dialysis.

9.4.10. **Platelets and Hemoglobin**

Platelet and hemoglobin measurements are not reliable with concurrent blood transfusions. Therefore, platelet values obtained from the day of a blood transfusion of platelets through 3 days after the transfusion will be excluded from all analyses. Hemoglobin values obtained from the day of a blood transfusion of either whole blood or packed red blood cells through 7 days after the transfusion will be excluded from all analyses. This rule will only be applied to postbaseline assessments of platelets and hemoglobin.
9.4.11. Assignment of CKD Stage

Chronic kidney disease (CKD) stage will be classified based on the National Kidney Foundation Chronic Kidney Disease Stage as follows:

Table 6: Stages of Chronic Kidney Disease

<table>
<thead>
<tr>
<th>Stage</th>
<th>Description</th>
<th>eGFR (mL/min/1.73 m²)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>Normal or High</td>
<td>≥90</td>
</tr>
<tr>
<td>2</td>
<td>Mildly Decreased(^a)</td>
<td>60-89</td>
</tr>
<tr>
<td>3a</td>
<td>Mildly to Moderately Decreased</td>
<td>45-59</td>
</tr>
<tr>
<td>3b</td>
<td>Moderately to Severely Decreased</td>
<td>30-45</td>
</tr>
<tr>
<td>4</td>
<td>Severely Decreased</td>
<td>15-29</td>
</tr>
<tr>
<td>5</td>
<td>Kidney failure</td>
<td>&lt; 15 (or dialysis)</td>
</tr>
</tbody>
</table>

\(^a\) Relative to young adult level

Note: In the absence of evidence of kidney damage, neither stage 1 nor 2 fulfill the criteria for CKD
9.5. Questionnaires and Patient Reported Outcomes

9.5.1. Pediatric FACIT-Fatigue

Below is a list of statements that other people with your illness have said are important. Please circle or mark one number per line to indicate your response as it applies to the past 7 days.

<table>
<thead>
<tr>
<th>Statement</th>
<th>None of the time</th>
<th>A little bit of the time</th>
<th>Some of the time</th>
<th>Most of the time</th>
<th>All of the time</th>
</tr>
</thead>
<tbody>
<tr>
<td>I feel tired</td>
<td>0</td>
<td>1</td>
<td>2</td>
<td>3</td>
<td>4</td>
</tr>
<tr>
<td>I have energy (or strength)</td>
<td>0</td>
<td>1</td>
<td>2</td>
<td>3</td>
<td>4</td>
</tr>
<tr>
<td>I could do my usual things at home</td>
<td>0</td>
<td>1</td>
<td>2</td>
<td>3</td>
<td>4</td>
</tr>
<tr>
<td>I had trouble <em>starting</em> things because I was too tired</td>
<td>0</td>
<td>1</td>
<td>2</td>
<td>3</td>
<td>4</td>
</tr>
<tr>
<td>I had trouble <em>finishing</em> things because I was too tired</td>
<td>0</td>
<td>1</td>
<td>2</td>
<td>3</td>
<td>4</td>
</tr>
<tr>
<td>I needed to sleep during the day</td>
<td>0</td>
<td>1</td>
<td>2</td>
<td>3</td>
<td>4</td>
</tr>
<tr>
<td>I got upset by being too tired to do things I wanted to do</td>
<td>0</td>
<td>1</td>
<td>2</td>
<td>3</td>
<td>4</td>
</tr>
<tr>
<td>Being tired made it hard for me to play or go out with my friends as much as I’d like</td>
<td>0</td>
<td>1</td>
<td>2</td>
<td>3</td>
<td>4</td>
</tr>
<tr>
<td>I needed help doing my usual things at home</td>
<td>0</td>
<td>1</td>
<td>2</td>
<td>3</td>
<td>4</td>
</tr>
<tr>
<td>I feel weak</td>
<td>0</td>
<td>1</td>
<td>2</td>
<td>3</td>
<td>4</td>
</tr>
<tr>
<td>I was too tired to eat</td>
<td>0</td>
<td>1</td>
<td>2</td>
<td>3</td>
<td>4</td>
</tr>
<tr>
<td>Being tired made me sad</td>
<td>0</td>
<td>1</td>
<td>2</td>
<td>3</td>
<td>4</td>
</tr>
<tr>
<td>Being tired made me mad (angry)</td>
<td>0</td>
<td>1</td>
<td>2</td>
<td>3</td>
<td>4</td>
</tr>
</tbody>
</table>
9.5.2. Resource Utilization Patient Questionnaire

Patient Questionnaire

RESOURCE UTILIZATION

1. Have you visited your health care provider for management of complications related to your aHUS within the past month?
   - No
   - Yes, specify number of health care provider visits in the past month: __________

2. Have you gone to an Emergency Room primarily for treatment of complications related to your aHUS in the past month?
   - No
   - Yes, specify number of emergency room visits in the past month: __________

3. Have you been admitted to the Hospital primarily for treatment of complications related to your aHUS in the past month?
   - No
   - Yes, specify number of hospital admissions in the past month: __________

4. Have you missed work or school as a result of complications or symptoms related to your aHUS within the past month?
   - No
   - Yes, specify number of work days missed in the past month: __________

GENERAL HEALTH

5. In general would you say your health is:
   - Excellent
   - Very good
   - Good
   - Fair
   - Poor

WORK STATUS

6. What is your current work status?
   - Work full time at a paid job
   - Work part time at a paid job (less than 30 hours per week)
   - Not in paid work force due to aHUS
   - Not working full time due to aHUS
   - Not working for pay for reasons unrelated to aHUS
   - Student
   - Retired
   - Other, specify: ___________________________________________
### 9.5.3. Patient-Reported AHUS Symptoms Questionnaire

<table>
<thead>
<tr>
<th>During the past week did you have any of the following symptoms?</th>
<th>Symptoms present?</th>
<th>If YES, how often did you have it?</th>
<th>If YES, how severe was it usually?</th>
<th>If YES, how much did it distress or bother you?</th>
</tr>
</thead>
<tbody>
<tr>
<td>Yellow discoloration of eyes and/or skin (jaundice)</td>
<td>□ No □ Yes</td>
<td>Rarely</td>
<td>Slight</td>
<td>Not at all</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Occasionally</td>
<td>Moderate</td>
<td>A little bit</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Frequently</td>
<td>Severe</td>
<td>Somewhat</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Almost constantly</td>
<td>Very Severe</td>
<td>Quite a bit</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>Very much</td>
</tr>
<tr>
<td>Chest pain</td>
<td>□ No □ Yes</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Shortness of breath</td>
<td>□ No □ Yes</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Headache</td>
<td>□ No □ Yes</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Confusion</td>
<td>□ No □ Yes</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Irritability</td>
<td>□ No □ Yes</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Anxiety</td>
<td>□ No □ Yes</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Weakness</td>
<td>□ No □ Yes</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Fatigue</td>
<td>□ No □ Yes</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Easy bruising/abdominal bleeding</td>
<td>□ No □ Yes</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Abdominal pain</td>
<td>□ No □ Yes</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Nausea/vomiting</td>
<td>□ No □ Yes</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Diarrhea</td>
<td>□ No □ Yes</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Swelling</td>
<td>□ No □ Yes</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Other, specify:</td>
<td>□ No □ Yes</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Other, specify:</td>
<td>□ No □ Yes</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Other, specify:</td>
<td>□ No □ Yes</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Other, specify:</td>
<td>□ No □ Yes</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Other, specify:</td>
<td>□ No □ Yes</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Other, specify:</td>
<td>□ No □ Yes</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Other, specify:</td>
<td>□ No □ Yes</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Date Completed: <strong><strong>/</strong></strong>/____</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>