

A trial to learn how safe AZD4512 is and how it works in the bodies of people with B-cell acute lymphoblastic leukemia (B-ALL)

Estado Reclutando	Tipo de Participantes Sujetos incapaces de otorgar consentimiento	Rangos de Edad Menores de 18 , Mayores de 64 , Adultos (18-64 años) , Adolescentes (12-17 años)
Género Ambos	Fases Fase I , Fase II	Participantes esperados 70
Resultados Sin resultados	Bajo nivel intervención No	Enfermedad rara Si
Cobertura geográfica Sin determinar	Ámbitos del ensayo seguridad, eficacia, farmacocinética, farmacodinámica, dosis	Terapia avanzada NO

Información

Identificador 2025-522372-93-00 (CTIS)	Cod. Protocolo ALLight D9891C00001
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Área terapéutica
Enfermedades [C] - Hematología [C15]

Enfermedad investigada
B-cell acute lymphoblastic leukemia (B-ALL)

Título Científico
A modular Phase I/II, open-label, multi-center study to evaluate the safety, tolerability, pharmacokinetics, immunogenicity, and preliminary efficacy of AZD4512 monotherapy or in combination with anticancer agent(s) in participants with Acute Lymphoblastic Leukemia

Justificación

No aportado

Objetivo Principal

Module 1 Dose Escalation: To assess the safety and tolerability of AZD4512 in participants with R/R B-ALL [Ph(-) and Ph(+)] as defined by NCCN guidelines

Module 1 Dose Escalation: To identify the MTD and/or doses of AZD4512 for subsequent evaluation in Module 2

Module 2 Dose Optimization: To evaluate the efficacy and determine the RP2D of AZD4512 in participants with R/R Ph(-) B-ALL based on NCCN response criteria

Module 2 Dose Optimization: To assess the safety and tolerability of AZD4512 in participants with R/R Ph(-) B-ALL

Variables de Evaluación Primaria

Module 1 Dose Escalation: Safety and Tolerability of AZD4512 -Assessment of DLT -Assessment of TEAEs/TRAES/SAEs -Interruptions, modifications, delays and discontinuations -Clinically significant changes from baseline

Module 1 Dose Escalation: Determine the MTD and/or doses to explore in Module 2

Module 2 Dose Optimization: Antitumour activity and determine the RP2D of AZD4512 -Response Rate: ORR (CR/CRh)

Module 2 Dose Optimization: Safety and Tolerability of AZD4512 -Assessment of TEAEs/TRAES/SAEs -Interruptions, modifications, delays and discontinuations -Clinically significant changes from baseline

Momentos temporales de evaluación primaria

No aportado

Objetivo Secundario

Module 1 Dose Escalation: To characterize the PK of AZD4512 as monotherapy

Module 1 Dose Escalation: To determine the immunogenicity of AZD4512 as monotherapy

Module 1 Dose Escalation: To evaluate the preliminary efficacy of AZD4512 as monotherapy in participants with R/R B-ALL [Ph(-) and Ph(+)] as defined by NCCN guidelines

Module 2 Dose Optimization: To evaluate the efficacy of AZD4512 based on NCCN response criteria

Module 2 Dose Optimization: To evaluate the impact of AZD4512 on MRD by NGS [central]

Module 2 Dose Optimization: To characterize the PK of AZD4512 as monotherapy

Module 2 Dose Optimization: To determine the immunogenicity of AZD4512 as monotherapy

Variables de Evaluación Secundaria

Module 1 Dose Escalation: Characterize AZD4512 PK as monotherapy

Module 1 Dose Escalation: Immunogenicity as monotherapy -ADA development

Module 1 Dose Escalation: Preliminary Antitumour Activity of AZD4512 -Response Rate: ORR (CR/CRh), CR and

CRc rate, TTR, DoR, EFS, OS, subsequent HSCT

Module 2 Dose Optimization: Antitumour Activity of AZD4512 -Response Rate: CR and CRc rate, TTR, DoR, EFS, OS, subsequent HSCT

Module 2 Dose Optimization: Effect of AZD4512 on MRD (NGS) -MRD-negative CR rate, CR/CRh (ORR), CRc (CR/CRi/CRh) rate

Module 2 Dose Optimization: PK of AZD4512 as monotherapy

Module 2 Dose Optimization: Immunogenicity as monotherapy -ADA development

Momentos temporales de evaluación secundaria

No aportado

Criterios de Inclusión

1. Age: 16 years old in Module 1 (US only: 18year) 12 years old in Module 2 2. Diagnosis: Diagnosis of B-ALL WHO (WHO-HAEM5) Participants must have relapsed or refractory B-ALL (relapsed defined as bone marrow blasts > 5% or reappearance of blasts in PB) - Module 1 (DE): Ph(-) B-ALL and Ph(+) B-ALL R/R - Backfill of Module 1 and Module 2 (DO): R/R Ph(-) B-ALL (BM blasts >5%) 3. Performance status (ECOG 2; KPS 50; LPS 50) 4. Peripheral lymphoblast count < 10,000/ μ L (may receive cytoreduction prior to C1D1 per protocol-specified criteria) 5. At least 2 prior therapies with refractoriness or relapse, or 1 prior therapy with refractoriness or relapse and no standard options available -Ph+ B-ALL (Module 1 DE only): intolerant to or have contraindications to TKI therapy or R/R disease despite treatment with at least 2 prior TKIs or at least one 3rd generation TKI 6. Prior DLI >4 weeks, prior cell therapy or autoHSCT >8 weeks, alloHSCT >12 weeks

Criterios de Exclusión

1. Burkitt lymphoma and leukemia 2. Isolated extramedullary disease; Active testicular or CNS (> CNS1) involvement 3. Unresolved non-heme toxicities Grade 2 (except alopecia, stable Grade 2 neuropathy, vitiligo, endocrine disorders controlled with therapy) 4. History of drug-induced non-infectious ILD/pneumonitis requiring oral or IV steroids or supplemental oxygen or where suspected ILD/pneumonitis cannot be ruled out by imaging at screening 5. Prior/concomitant therapy -Cytotoxic treatment within 14 days (except ALL maintenance medications or cytoreduction) -Biologic (immuno-oncology) treatment within 28 days or 5 half-lives (whichever is shorter) -Non-CNS radiation within 2 weeks & CNS radiation within 4 weeks -Medications known to prolong QTc and/or associated with Torsades de Pointes within 5 half-lives -Strong inhibitors of CYP 3A4 within 14 days or 5 half-lives (whichever is longer) -Investigational agents or study interventions in the last 30 days or 5 half-lives prior to the first dose of AZD4512 whichever is longer. If the investigational product is an agent to treat B-ALL and meets the modality criteria, then a specific washout period must be adhered to instead.

Calendario

(Última actualización: 10/06/2026)

Autorización 24/11/2025	Inicio de Ensayo 13/01/2026	Inclusión Primer Paciente 19/01/2026	Interrumpido No aportado	Reiniciado No aportado
Fin de reclutamiento No aportado	Fin prematuro (España) No aportado	Fin prematuro (Global) No aportado	Fin del ensayo en España No aportado	Fin del ensayo global No aportado

Promotor

AstraZeneca AB Sweden

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Contact Person

AstraZeneca Clinical Study Information Center

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Monetary support: N/A

Tipo de promotor: Comercial

Ceim

CEIm de Cantabria

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Centros

Hospital Germans Trias I Pujol

Badalona

BARCELONA

Servicio de Hematología

No iniciado (---/---/----)

Hospital Universitari Vall D Hebron

Barcelona

BARCELONA

Servicio de Hematología

Activo (13/01/2026)

Hospital Universitario De Salamanca

Salamanca

SALAMANCA

Servicio de Hematología

Activo (22/01/2026)

Hospital Universitario Marques De Valdecilla

Santander

CANTABRIA

Servicio de Hematología

Activo (05/02/2026)

Medicamentos

AZD4512
SOLUTION FOR INJECTION/INFUSION
IV INFUSION

Principios Activos: N/A

Experimental

Sin resultados

A trial to learn how safe AZD4512 is and how it works in the bodies of people with B-cell acute lymphoblastic leukemia (B-ALL)

State Recruiting	Type of participants Incapable subjects of giving consent	Age Ranges Less than 18 , Older than 64 , Adults (18-64 years) , Teens (12-17 years)
Gender Both	Phases Phase I , Phase II	Expected Participants 70
Results No results	Low level of intervention No	Rare disease Yes
Geographic coverage Undetermined	Areas of the study safety, effectiveness, pharmacokinetics, pharmacodynamics, dose	Advanced therapy NO

Information

Identifier

2025-522372-93-00 (CTIS)

Protocol Code

ALLight D9891C00001

Therapeutic area

Diseases [C] - Hemic and Lymphatic Diseases [C15]

Investigated Disease

B-cell acute lymphoblastic leukemia (B-ALL)

Scientific Title

A modular Phase I/II, open-label, multi-center study to evaluate the safety, tolerability, pharmacokinetics, immunogenicity, and preliminary efficacy of AZD4512 monotherapy or in combination with anticancer agent(s) in participants with Acute Lymphoblastic Leukemia

Rationale

Not provided

Main Objective

Module 1 Dose Escalation: To assess the safety and tolerability of AZD4512 in participants with R/R B-ALL [Ph(-) and Ph(+)] as defined by NCCN guidelines

Module 1 Dose Escalation: To identify the MTD and/or doses of AZD4512 for subsequent evaluation in Module 2

Module 2 Dose Optimization: To evaluate the efficacy and determine the RP2D of AZD4512 in participants with R/R Ph(-) B-ALL based on NCCN response criteria

Module 2 Dose Optimization: To assess the safety and tolerability of AZD4512 in participants with R/R Ph(-) B-ALL

Primary Endpoints

Module 1 Dose Escalation: Safety and Tolerability of AZD4512 -Assessment of DLT -Assessment of TEAEs/TRAES/SAEs -Interruptions, modifications, delays and discontinuations -Clinically significant changes from baseline

Module 1 Dose Escalation: Determine the MTD and/or doses to explore in Module 2

Module 2 Dose Optimization: Antitumour activity and determine the RP2D of AZD4512 -Response Rate: ORR (CR/CRh)

Module 2 Dose Optimization: Safety and Tolerability of AZD4512 -Assessment of TEAEs/TRAES/SAEs -Interruptions, modifications, delays and discontinuations -Clinically significant changes from baseline

Temporary moments of secondary assessment

Not provided

Secondary Objective

Module 1 Dose Escalation: To characterize the PK of AZD4512 as monotherapy

Module 1 Dose Escalation: To determine the immunogenicity of AZD4512 as monotherapy

Module 1 Dose Escalation: To evaluate the preliminary efficacy of AZD4512 as monotherapy in participants with R/R B-ALL [Ph(-) and Ph(+)] as defined by NCCN guidelines

Module 2 Dose Optimization: To evaluate the efficacy of AZD4512 based on NCCN response criteria

Module 2 Dose Optimization: To evaluate the impact of AZD4512 on MRD by NGS [central]

Module 2 Dose Optimization: To characterize the PK of AZD4512 as monotherapy

Module 2 Dose Optimization: To determine the immunogenicity of AZD4512 as monotherapy

Secondary Endpoints

Module 1 Dose Escalation: Characterize AZD4512 PK as monotherapy

Module 1 Dose Escalation: Immunogenicity as monotherapy -ADA development

Module 1 Dose Escalation: Preliminary Antitumour Activity of AZD4512 -Response Rate: ORR (CR/CRh), CR and

CRc rate, TTR, DoR, EFS, OS, subsequent HSCT

Module 2 Dose Optimization: Antitumour Activity of AZD4512 -Response Rate: CR and CRc rate, TTR, DoR, EFS, OS, subsequent HSCT

Module 2 Dose Optimization: Effect of AZD4512 on MRD (NGS) -MRD-negative CR rate, CR/CRh (ORR), CRc (CR/CRi/CRh) rate

Module 2 Dose Optimization: PK of AZD4512 as monotherapy

Module 2 Dose Optimization: Immunogenicity as monotherapy -ADA development

Temporary moments of secondary assessment

Not provided

Inclusion criteria

1. Age: 16 years old in Module 1 (US only: 18year) 12 years old in Module 2 2. Diagnosis: Diagnosis of B-ALL WHO (WHO-HAEM5) Participants must have relapsed or refractory B-ALL (relapsed defined as bone marrow blasts > 5% or reappearance of blasts in PB) - Module 1 (DE): Ph(-) B-ALL and Ph(+) B-ALL R/R - Backfill of Module 1 and Module 2 (DO): R/R Ph(-) B-ALL (BM blasts >5%) 3. Performance status (ECOG 2; KPS 50; LPS 50) 4. Peripheral lymphoblast count < 10,000/ μ L (may receive cytoreduction prior to C1D1 per protocol-specified criteria) 5. At least 2 prior therapies with refractoriness or relapse, or 1 prior therapy with refractoriness or relapse and no standard options available -Ph+ B-ALL (Module 1 DE only): intolerant to or have contraindications to TKI therapy or R/R disease despite treatment with at least 2 prior TKIs or at least one 3rd generation TKI 6. Prior DLI >4 weeks, prior cell therapy or autoHSCT >8 weeks, alloHSCT >12 weeks

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Calendar

(Last Update: 10/06/2026)

Authorization 24/11/2025	Start of Trial 13/01/2026	First patient inclusion 19/01/2026	Halted Not aported	Restarted Not aported
End of recruitment Not aported	Premature end (Spain) Not aported	Premature End (Global) Not aported	Trial end (Spain) Not aported	Trial end (Global) Not aported

Sponsor

AstraZeneca AB Sweden

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Contact Person

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Monetary support: N/A

Sponsor type: Commercial

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Sites

not initialized (---/---/----)	Hospital Germans Trias I Pujol Badalona BARCELONA Servicio de Hematologia
Active (13/01/2026)	Hospital Universitari Vall D Hebron Barcelona BARCELONA Servicio de Hematologia
Active (22/01/2026)	Hospital Universitario De Salamanca Salamanca SALAMANCA Servicio de Hematologia
Active (05/02/2026)	Hospital Universitario Marques De Valdecilla Santander CANTABRIA Servicio de Hematologia

Medication

AZD4512 SOLUTION FOR INJECTION/INFUSION IV INFUSION
- Active Principles: N/A Experimental

No results