

Masitinib Significantly Decreases the Rate of Asthma Exacerbations in Patients with Severe Asthma Uncontrolled by Oral Corticosteroids: A phase 3 Multicenter Study

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On behalf of the AB07015 Study Group

Conflict of interest disclosure



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- ☐ I have no real or perceived conflicts of interest that relate to this presentation.
- ✓ I have the following real or perceived conflicts of interest that relate to this presentation:

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Other support / potential conflict of interest:



Masitinib is a first-in-class oral tyrosine kinase inhibitor in severe asthma

- Masitinib selectively targets mast cell activity (c-Kit, LYN, FYN) and is also a potent inhibitor of PDGFR
- Strong scientific rationale to target mast cells in severe asthma
- PDGFR signaling associated with airway remodeling in severe asthma
- Masitinib activity demonstrated via preclinical mouse models of asthma
 - Significant decrease of airway hyper-responsiveness
 - Significant decrease of eosinophils recruitment
- Clinical proof-of-concept in cat [Lee-Fowler, 2012] and human [Humbert, 2009] studies



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Study AB07015 evaluated masitinib 6.0 mg/kg/day in severe asthma uncontrolled by OCS with no restriction on baseline eosinophil level

- Randomized (2:1), double-blinded, placebo-controlled.
- Patient with severe asthma, uncontrolled by OCS (≥7.5 mg/d), both high (≥150 cells/μL) and low (<150 cells/μL) eosinophils
- Timing: 2-week run-in (blinded placebo) → 36-week treatment period [W0–W36]
 → possible blinded extension
- Primary endpoint: Reduction of annualized severe asthma exacerbation rate for overall exposure
- If significant, sequential analysis in pts with severe asthma and eosinophil count
 ≥150 cells/μL (fixed hierarchical procedure with 5% α for each analysis)

Study AB07015 Primary Endpoint Results



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Masitinib significantly decreased the rate of severe asthma exacerbations in patients with severe asthma uncontrolled by OCS, regardless of eosinophil level

Primary Analysis (Severe Asthma) Annualized severe asthma exacerbation rate					Sequential Analysis (Severe Asthma with High Eosinophil) Annualized severe asthma exacerbation rate						
Anr	Exp			_	e P-value	Ani	nualize Exp		RR [95%CI]		re P-value
MAS (240)	1.14	0.34	0.65	35%	0.0103	MAS (181)	1.10	0.34	0.62	200/	0.015/
PBO (115)	1.15	0.48	[0.47, 0.90]			PBO (87)	1.12	0.51	[0.42, 0.91]	38%	0.0156
Exp: Exposure in years. RR: rate ratio. MAS: Masitinib 6.0 mg/kg/d. PBO: Placebo.											

- Average exposure (approx. 60 weeks) was well-balanced across treatment-arms
- Sensitivity analysis on ITT population was similarly significant (-33%, p-value=0.0156)

Sensitivity Analysis on Primary Endpoint



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Benefit of masitinib was greatest in patients who had higher cumulated use of OCS (indicative of more severe asthma that is harder to control)

Sensitivity Analysis (Severe Asthma)				Sensitivity Analysis (Severe Asthma with High Eosinophil)							
Annualized severe asthma exacerbation rate				Annualized severe asthma exacerbation rate							
Cumulative OCS >500 mg				Cumulative OCS >500 mg							
	Exp	Rate	RR [95%CI]	Reduction	P-value	_	Exp	Rate	RR [95%CI]	Reduction	P-value
MAS (161)	1.15	0.34	0.59	47.0/	0.0092	MAS (127)	1.12	0.32	0.51	49%	0.0049
PBO (82)	1.20	0.55	[0.39, 0.88]	41%		PBO (60)	1.16	0.60	[0.32, 0.82]		
Cumulative OCS >1000 mg					Cumulative OCS >1000 mg						
		Rate	RR [95%CI]	Reduction	P-value	_		Rate	RR [95%CI]	Reduction	P-value
MAS (120)	1.16	0.26	0.49	51%	0.0060	MAS (92)	1.11	0.22	0.29	710/	0.0003
PBO (66)	1.27	0.53	[0.29, 0.82]			PBO (46)	1.27	0.55	[0.15, 0.57]	71%	0.0003
Exp: Exposure in years. RR: rate ratio. MAS: Masitinib 6.0 mg/kg/d. PBO: Placebo. OCS oral corticosteroid.											



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Safety was consistent with the known profile for masitinib

Summary of Adverse Events (AE) - Safety Population (n = 404)

	Masitinib % (n)	Placebo % (n)	Difference
At least one AE	83.4% (226/271)	82.0% (109/133)	+1.4%
At least one severe AE	48.0% (130/271)	45.9% (61/133)	+2.1%
At least one serious AE (non-fatal)	17.7% (48/271)	16.5% (22/133)	+1.2%



No new safety signals were observed



Study AB07015 demonstrated efficacy in a difficult to treat population

- Study AB07015 population is distinct from other asthma trials
 - Patients dependent on OCS (100% receiving high dose OCS therapy) and no weaning
 - Patients were treated irrespective of baseline eosinophil count
 - Evaluated over a long period of time (approx. 60 weeks)
- Significant reduction in severe asthma exacerbation rate
 - -35% reduction in primary analysis population irrespective of baseline eosinophil level
 - -38% reduction in subgroup with baseline eosinophil level ≥150 cells/μL
 - Greatest benefit (-41% to -71%) for patients who had the most severe asthma
- Masitinib may therefore provide a new treatment option for severe asthma uncontrolled by OCS
 - Biologic-ineligible patients (e.g. eosinophil count of ≤300 cells/μL)
 - Patients in failure to biologics

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