

**MTS1 : Myopia Treatment Study 1**  
**Month 9 Call**  
**9-month call Follow-Up Form**

Patient ID: **M01-004-2299**

**SECTION: PHONE CALL INFORMATION**

**(VisitInformation\_d\_05)**

**CONTACT INFORMATION**

**1. Name of person conducting contact:** tblMTS1VisitInfo.dbo.PhoneCallUserID

21 - Shelly T. Mares

BM1: tblMTS1VisitInfo.dbo.ColumnName\_Property\_Not\_Set\_In\_Template

**2. Contact date:** tblMTS1VisitInfo.dbo.VisitDt tblMTS1VisitInfo.dbo.VisitMiss

19 Jan 2018

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Missed

If Missed, reason: tblMTS1VisitInfo.dbo.VisitMissReason

If Other, describe: tblMTS1VisitInfo.dbo.VisitMissReasonDs

**OUT OF TARGET WINDOW**

tblMTS1VisitInfo.dbo.OutOfWin **Contact was completed out of target window**

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**1. Reason contact was completed out of target window:** tblMTS1VisitInfo.dbo.OutOfWinReason

Poor outcome

1a. If Other, describe: tblMTS1VisitInfo.dbo.OutOfWinReasonDs

**SECTION: PHONE CALL FORM**

**(MTS1FUCall\_a\_01)**

**PHONE CALL INFORMATION**

**1. Person spoken with:** tblMTS1FUCall.dbo.MTS1PersSpokenWith

Father

1a. If Other, describe: tblMTS1FUCall.dbo.MTS1PersSpokenWithDs

**2. Has the participant been taking the prescribed study medication?** tblMTS1FUCall.dbo.TakingMed

Yes

2a. If No, why not (consult with investigator if necessary)? tblMTS1FUCall.dbo.TakingMedDs

**3. Has the participant experienced any side effects or other adverse effects since the last visit?** tblMTS1FUCall.dbo.SideEffects

No

3a. If Yes, describe: tblMTS1FUCall.dbo.SideEffectsDs

**Complete an Adverse Event form if the event or medical problem meets any of the following criteria and consult with investigator if necessary.**

An Adverse Event form should be completed if the participant reports any of the following new events since the last visit:

1. If the participant has experienced any new ocular side effects of treatment including lid/conjunctival irritation, light sensitivity, or near blur and/or reading difficulty.
2. If the participant has experienced any systemic side effects of treatment presenting within one hour following administration of study medication, including dry skin/mouth, tachycardia, fever, flushing, irritability, mental confusion, constipation, aggravation of asthma, or seizures.
3. If the participant has experienced any medical problem meeting criteria for a serious adverse event as follows: A serious adverse event is any untoward occurrence that:
  - Results in death.
  - Is life-threatening; (a non-life-threatening event which, had it been more severe, might have become life-threatening, is not necessarily considered a serious adverse event).
  - Requires inpatient hospitalization or prolongation of existing hospitalization.
  - Results in persistent or significant disability/incapacity or substantial disruption of the ability to conduct normal life functions (sight-threatening).
  - Is considered a significant medical event by the investigator based on medical judgment (e.g., may jeopardize the participant or may require

medical/surgical intervention to prevent one of the outcomes listed above).

**COMMENTS**

tblMTS1FUCall.dbo.CallCmts