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CUI DEVICES

date 10/14/2020

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MODEL: CLS0282MA-L152 | DESCRIPTION: SPEAKER

FEATURES

- 28 mm
- round frame
- 0.5 W
- 8 Ω
- neodymium magnet
- mylar cone
- wire leads





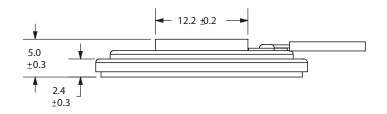
SPECIFICATIONS

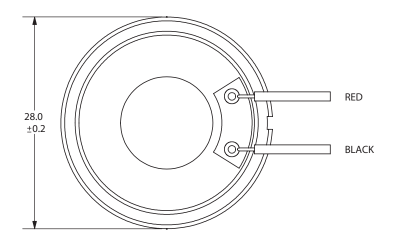
parameter	conditions/description	min	typ	max	units
input power			0.5	1.0	W
impedance	at 1.5 kHz, 1.0 V	6.8	8	9.2	Ω
resonant frequency (Fo)	at 1.0 V	400	550	700	Hz
frequency response		Fo		10,000	Hz
sound pressure level	at 0.5 W, 50 cm, avg at 0.8, 1.0, 1.2, 1.5 kHz	82	85	88	dB
buzz, rattle, etc.	must be normal at sine wave			2.0	V
dimensions	Ø28 x 5				mm
magnet	Nd-Fe-B				
cone material	mylar				
terminal	wire leads				
weight			5.8		g
operating temperature		-25		60	°C
hand soldering	for maximum 3 seconds	340	350	360	°C
RoHS	yes				

MECHANICAL DRAWING

units: mm

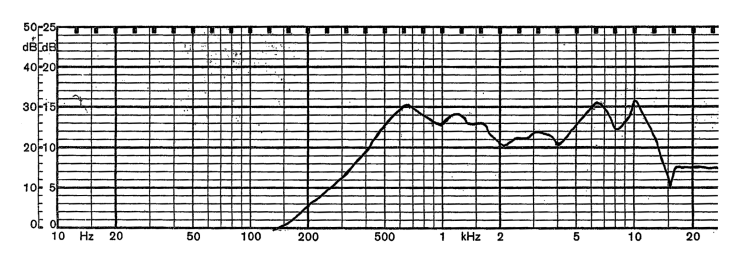
wire: UL1007 28 AWG length: 152 mm





FREQUENCY RESPONSE CURVE

parameter	conditions/description
potentiometer range	50 dB
rectifier	RMS
lower limit frequency	20 Hz
wr. speed	100 mm/sec
zero level	60 dB



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CUI Devices | MODEL: CLS0282MA-L152 | DESCRIPTION: SPEAKER date 10/14/2020 | page 3 of 3

REVISION HISTORY

rev.	description	date
1.0	initial release	07/03/2007
1.01	brand update	10/14/2020

The revision history provided is for informational purposes only and is believed to be accurate.

CUI DEVICES

CUI Devices offers a one (1) year limited warranty. Complete warranty information is listed on our website.

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CUI Devices products are not authorized or warranted for use as critical components in equipment that requires an extremely high level of reliability. A critical component is any component of a life support device or system whose failure to perform can be reasonably expected to cause the failure of the life support device or system, or to affect its safety or effectiveness.