

MODEL: CPS-4242-100T | **DESCRIPTION:** SIREN**FEATURES**

- siren tone
- internally driven
- through hole

**SPECIFICATIONS**

parameter	conditions/description	min	typ	max	units
rated voltage			12		Vdc
operating voltage		6		15	Vdc
current consumption	at rated voltage			200	mA
rated frequency		1,500		4,000	Hz
sound pressure level	at 30 cm, rated voltage	100			dB
tone	siren, at rated voltage				
dimensions	Ø42.5 x 42.0				mm
weight				60.8	g
material	ABS (UL94 1/16" HB)				
terminal	pins (tin plating)				
operating temperature		-30		85	°C
storage temperature		-40		95	°C
washable	no				
RoHS	yes				

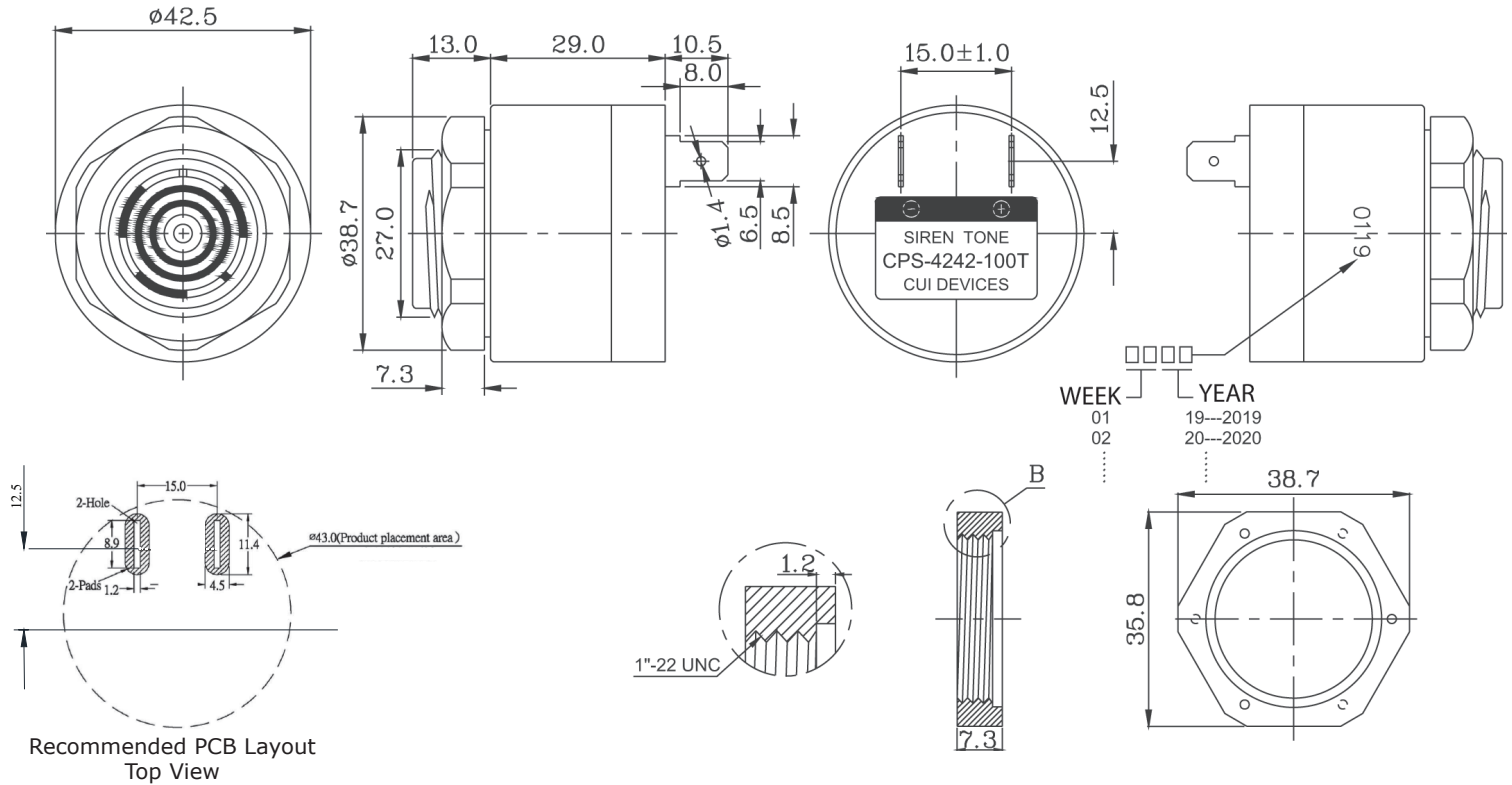
Notes: 1. All specifications measured at 5~35°C, humidity at 45~85%, under 86~106 kPa pressure, unless otherwise noted.

SOLDERABILITY

parameter	conditions/description	min	typ	max	units
hand soldering	maximum 3 seconds	330	350	370	°C

MECHANICAL DRAWING

units: mm

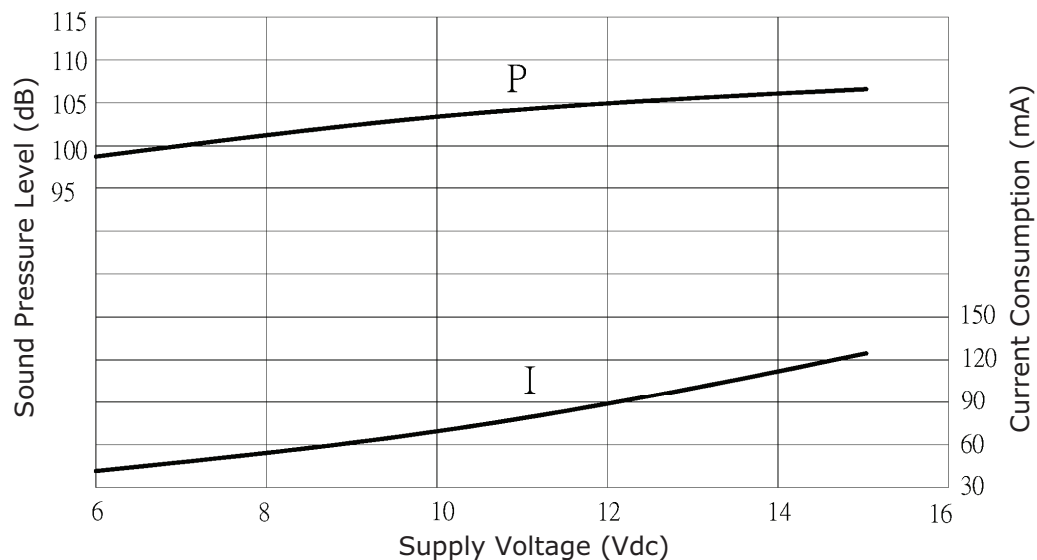
tolerance: ± 0.5 mm

PERFORMANCE CURVES

P: Voltage vs. Sound Pressure Level

I: Voltage vs. Current Consumption

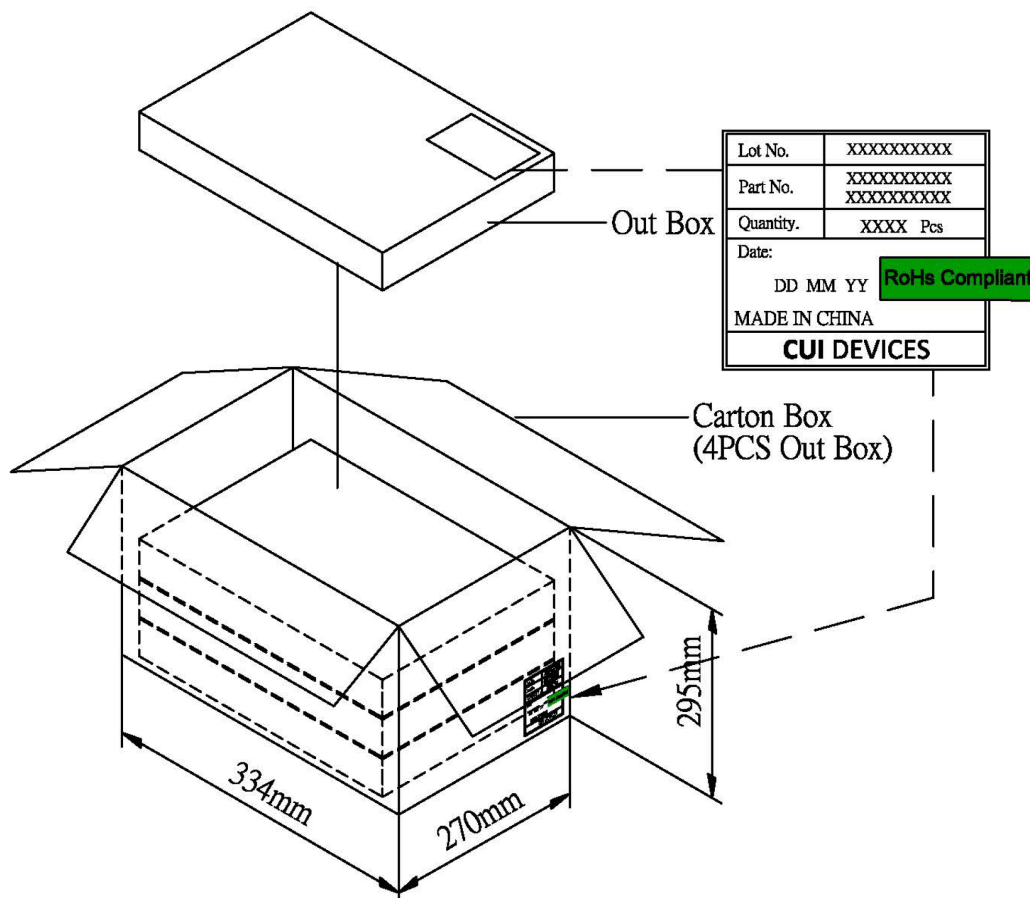
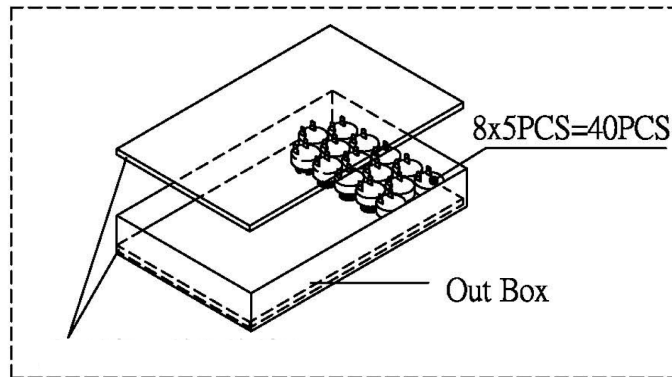
at 30 cm



PACKAGING

units: mm

Carton Size: 334 x 270 x 295 mm
Carton QTY: 160 pcs per carton



REVISION HISTORY

rev.	description	date
1.0	initial release	01/22/2020

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

CUI DEVICES

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