



IN PARTNERSHIP WITH



WHO WE SERVE

Children living with Alopecia
and their families



OUR VISION

Changing the emphasis
from growing hair to
growing confidence,
building self-esteem,
providing support and
raising awareness of
Alopecia.



OUR MISSION

To help any child in need
who is living with hair loss
due to all forms of Alopecia.

How Alopecia
IMPACT
patients



We show them that everyone is different and that different is normal. They are perfect as they are, and if they want to change something they can control, change it. If you can't. Own it. People like you because of who you are, not what you look like, and if that is true with anyone, you need to find others to fill your friend list.

HOW WE MAKE AN IMPACT

The Children's Alopecia Project (CAP) is a nonprofit organization dedicated to supporting children with alopecia by building self-esteem, confidence, and community.

CAP provides programs, resources, and events designed to help kids with alopecia feel connected, empowered, and understood.

The Children's Alopecia Project is committed to helping kids with alopecia feel seen, supported, and confident just as they are.



AT A GLANCE HOW WE PROVIDE SUPPORT

Alopeciapalooza, Camps, Get-togethers, support groups, CAP creates opportunities for children and families to connect with others living with alopecia. Alopecia awareness and education through initiatives like the CAP2U Speaking Tour, helping schools and communities better understand alopecia and support inclusivity.

GET SOCIAL WITH US



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SCAN TO LEARN MORE ABOUT NEVUS OUTREACH



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