Pausing for The GFPD Op-Ed 2023 Templates

Instructions:
1. Locate an Op-Ed template from the choices below that best matches your family (bereaved parent or not, adult patient, type of peroxisomal disorder, etc.)
2. Copy and paste the template to your own document.
3. Customize the template to fit you and your family.
4. Submit to your local newspaper. This is usually through an online form on their website.
5. Let the GFPD know when your Op-Ed runs in the newspaper by contacting Katie at katie@thegfpd.org

Glossary of Terms:
PBD-ZSD = Peroxisomal Biogenesis Disorder – Zellweger Syndrome Disorder
DBPD = D-Bifunctional Protein Deficiency
ACOX = Acyl-CoA Oxidase Deficiency
AMACRD = 2-Methylacyl-CoA racemase Deficiency

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To the Editor:

October 5th is a special day for our family as we celebrate Pause for The GFPD, the annual awareness day of The Global Foundation for Peroxisomal Disorders (GFPD). I invite the NEWSPAPER NAME readership to join our family and PAUSE for families impacted by peroxisomal disorders in more than 40 countries around the world.

OUR/MY DAUGHTER/SON, NAME, lived for _______ months/years with a peroxisomal biogenesis disorder in the Zellweger spectrum (PBD-ZSD). CHILDS NAME is one of fewer than 1,000 children worldwide affected with PBD-ZSD. TELL 1-3 SENTENCES ABOUT YOUR CHILD.

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On October 5th, we are celebrating all that the GFPD has accomplished in the last 13 years in moving towards a cure for peroxisomal disorders. Most importantly, Pause for The GFPD celebrates the beautiful children affected by this rare, terminal condition. We invite you to help the GFPD – and our global community of patients, families, caregivers, scientists, and medical professionals – raise awareness about the need for treatments and cures for this rare, genetic, and terminal disease.

Raising public awareness makes a difference. It gives families hope and can lead to new, life-saving treatments. Taking the time to participate in Pause for The GFPD also helps those of us who have lost a child to know our child is still loved and remembered. I encourage everyone reading this to get involved by visiting the GFPD website: http://www.thegfpd.org, or find us on Facebook, Instagram, or Twitter @thegfpd to learn about Pause for The GFPD celebrations in your area, ways you can help make a difference, and to learn more about the groundbreaking work the GFPD is doing around the world.

Sincerely,
YOUR NAME
YOUR CITY, YOUR STATE
Template for Parent(s) of Child(ren) Living with a Peroxisomal Biogenesis Disorder in the Zellweger Spectrum (PBD-ZSD)

To the Editor:

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OUR/MY DAUGHTER/SON, NAME, is HOW MANY years old and is one of fewer than 1,000 children living worldwide with a peroxisomal biogenesis disorder in the Zellweger spectrum (PBD-ZSD). TELL 1-3 SENTENCES ABOUT YOUR CHILD. PBD-ZSD is a rare, genetic, condition affecting multiple organ systems in the body, and is generally fatal in childhood.

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Sincerely,
YOUR NAME
YOUR CITY, YOUR STATE
Template for Adult Living with a
Peroxisomal Biogenesis Disorder in the Zellweger Spectrum (PBD-ZSD)

To the Editor:

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I am one of fewer than 1,000 individuals living worldwide with a peroxisomal biogenesis disorder in the Zellweger spectrum (PBD-ZSD). TELL 1-3 SENTENCES ABOUT YOURSELF AND HOW PBD IMPACTS YOUR LIFE. PBD-ZSD is a rare, genetic, condition affecting multiple organ systems in the body, and is generally fatal in childhood.

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Sincerely,
YOUR NAME
YOUR CITY, YOUR STATE
Template for Parent(s) of a Child(ren) Who Died From D-Bifunctional Protein Deficiency (DBPD)

To the Editor:

October 5th is a special day for our family as we celebrate Pause for The GFPD, the annual awareness day of The Global Foundation for Peroxisomal Disorders (GFPD). I invite the NEWSPAPER NAME readership to join our family and PAUSE for families impacted by peroxisomal disorders in more than 40 countries around the world.

OUR/MY DAUGHTER/SON, NAME, lived for ______ months/years with a peroxisomal disorder called, d-bifunctional protein deficiency (DBPD). TELL 1-3 SENTENCES ABOUT YOUR CHILD.

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Sincerely,
YOUR NAME
YOUR CITY, YOUR STATE
Template for Parent(s) of Child(ren) Living with D-Bifunctional Protein Deficiency (DBPD)

To the Editor:

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OUR/MY DAUGHTER/SON, NAME, is HOW MANY years old and is one of only a few children living worldwide with a peroxisomal disorder called, d-bifunctional protein deficiency (DBPD). TELL 1 - 3 SENTENCES ABOUT YOUR CHILD. DBPD is a rare, genetic, condition affecting multiple organ systems in the body, and is generally fatal in childhood.

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Sincerely,
YOUR NAME
YOUR CITY, YOUR STATE
Template for Adult(s) Living with D-Bifunctional Protein Deficiency (DBPD)

To the Editor:

October 5th is a special day for me and my family as we celebrate Pause for The GFPD, the annual awareness day of The Global Foundation for Peroxisomal Disorders (GFPD). I invite the NEWSPAPER NAME readership to join me and PAUSE for the families like mine, in more than 40 countries around the world, who are impacted by peroxisomal disorders.

I am one of only a few in the world who are living with a peroxisomal disorder called d-bifunctional protein deficiency (DBPD). TELL 1-3 SENTENCES ABOUT YOURSELF AND HOW DBPD IMPACTS YOUR LIFE. DBPD is a rare, genetic, condition affecting multiple organ systems in the body, and is generally fatal in childhood.

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Sincerely,
YOUR NAME
YOUR CITY, YOUR STATE
Template for Parent(s) of a Child(ren) Who Died From
Acyl-CoA Oxidase Deficiency (ACOX)

To the Editor:

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OUR/MY DAUGHTER/SON, NAME, lived for ______ months/years with a peroxisomal disorder called, Acyl-CoA Oxidase Deficiency (ACOX). TELL 1-3 SENTENCES ABOUT YOUR CHILD.

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Template for Parent(s) of Child(ren) Living with
Acyl-CoA Oxidase Deficiency (ACOX)

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OUR/MY DAUGHTER/SON, NAME, is HOW MANY years old and is one of only a few children living worldwide with a peroxisomal disorder called, Acyl-CoA Oxidase Deficiency (ACOX). TELL 1 -3 SENTENCES ABOUT YOUR CHILD. ACOX is a rare, genetic, condition affecting multiple organ systems in the body, and is generally fatal in childhood.

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I am one of only a few in the world who are living with a peroxisomal disorder called Acyl-CoA Oxidase Deficiency (ACOX). TELL 1-3 SENTENCES ABOUT YOURSELF AND HOW ACOX IMPACTS YOUR LIFE. ACOX is a rare, genetic, condition affecting multiple organ systems in the body, and is generally fatal in childhood.

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OUR/MY DAUGHTER/SON, NAME, lived for _____ months/years with a peroxisomal disorder called, 2-Methylacyl-CoA racemase Deficiency (AMACRD). TELL 1-3 SENTENCES ABOUT YOUR CHILD.

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Sincerely,
YOUR NAME
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Template for Parent(s) of Child(ren) Living with 2-Methylacyl-CoA racemase Deficiency (AMACRD)

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I am one of only a few in the world who are living with a peroxisomal disorder called 2-Methylacyl-CoA racemase Deficiency (AMACRD). TELL 1-3 SENTENCES ABOUT YOURSELF AND HOW AMACR Deficiency impacts YOUR LIFE. AMACR Deficiency is a rare, genetic, condition.

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