HEADACHE & MIGRAINE DISEASE LANGUAGE & IMAGE GUIDE 2018/19





This document represents the current consensus of the headache, migraine and cluster disease patient community. Language is constantly evolving and we will update this guide over time so that it continues to accurately represent how people living with headache diseases want our conditions described and portrayed.

Version 1.1 (July 2019)

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Declaration of Rights for People with Headache Diseases



We, the people with headache, migraine and cluster disease, deserve respect and empathy for the serious and sometimes debilitating medical conditions with which we live. We will not be disempowered, minimized or stigmatized by a society that does not understand the complexity and seriousness of our disease. We demand research, treatments and societal support that is proportional to the impact of our disease. This means an increase in funding for research, better insurance policies that make it easier for us to access our treatments and more visibility in the public space.

We are working to educate people on why the terms "patient" and "sufferer" can be problematic. Sufferers, implies defeat, and we are patients only when interacting with medical professionals who manage our diseases – we are people living with migraine, cluster or headache disease 24/7.

We will fight to have our struggle against these painful and disabling diseases honored. We, to the best of our ability, will make headache diseases visible.

Recommendations For All People



1. Honor and believe the struggle of people living with headache, migraine and cluster disease to cope with their pain and many other symptoms.

2. Educate and work to correct the culture that penalizes people with severe headache, migraine and cluster disease.

3. Learn about these diseases and understand the systemic nature of them.

Recommendations For People With Headache Diseases

1. When you can, reject the stigma of headache, migraine and cluster in yourself, and confront it in others. This takes work and time to understand the messages we have received about living with a chronic disease.



3. When you can, participate in awareness and advocacy efforts. Seek out policy makers and advocate for understanding, research and treatment.

Rights Of People With Headache Diseases

1. To live a life of dignity, despite whatever limitations may be caused by our disease.

- 2. To not be blamed or dismissed for having a headache disease when all we want is effective treatment.
- 3. To gain access to quality medical treatment and social service provision not limited by external forces, financial or otherwise.
- 4. To advance robust scientific research that addresses the impact of headache disease and chronic pain on people living with these diseases, their caregivers and families and society.

Why Disease Language & Images Matter

Language plays a vital role in how we understand the world. Too often, language can be used to disparage and belittle people living with disease, labeling them as helpless or as victims. This ultimately creates shame and stigma, and illegitimately justifies the notion that people with diseases should be ignored

or feared. We see this prominently played out in the headache disease area. Language used by medical experts, the media, the public and stakeholders of our own community can greatly impact how people living with headache diseases are

perceived and treated. This, in turn, affects the resources society makes available to support our pain and disability.

Likewise, the images associated with headache, migraine and cluster diseases often do not accurately depict the full range of symptoms connected to these diseases and minimizes the severity of impact on daily lives. Images used by media or depicted in popular culture often overlook the diversity of races, ages and genders that are impacted by headache diseases. This adds to the mislabeling, misunderstanding, minimization and stigmatization of the individuals living with these diseases. not intend this guide to be used to badger or shame anyone for the language they use. We hope this document will be used as a guiding light to help steer the way to more compassion and understanding for those living with migraine, cluster and other headache diseases.

The purpose of this guide is to help all people who communicate about headache diseases to do so in a more accurate, informed and empathetic manner that treats these conditions and the people living with them with the seriousness and care they deserve.

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Language is constantly evolving, and we will update this guide over time so that it reflects the most updated discussions of the headache patient advocacy community. We do Dr. William B. Young led and published a 2011 study, Naming Migraine and Those Who Have It¹, that convened a diverse consensus panel and analyzed its responses to a series of questions related to the language used for migraine. This guide

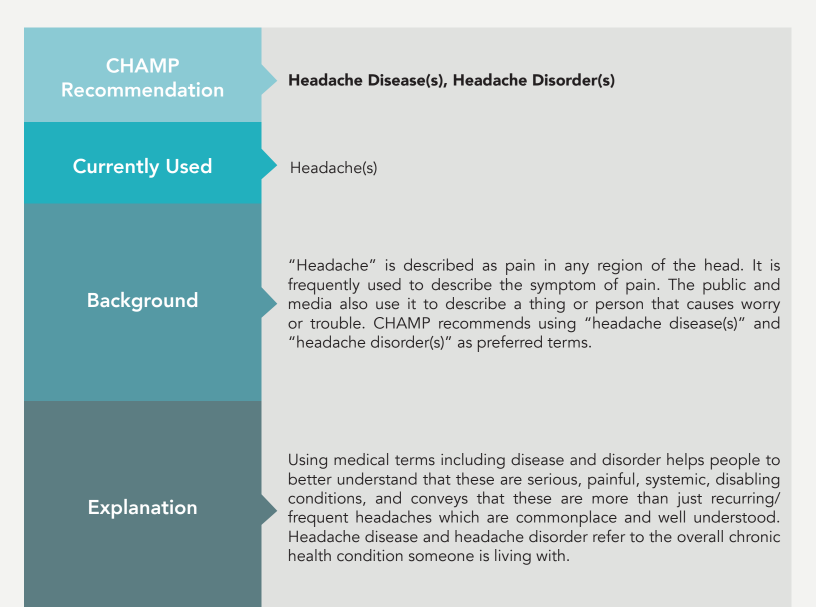
builds upon that important work.

The Coalition For Headache And Migraine Patients (CHAMP) acknowledges and thanks Dr. Young and his colleagues for launching this conversation. We, the headache, migraine and cluster patient advocacy community, are honored to use this important work as a foundation for our own.



Language Guide

Headache Disease – Overall



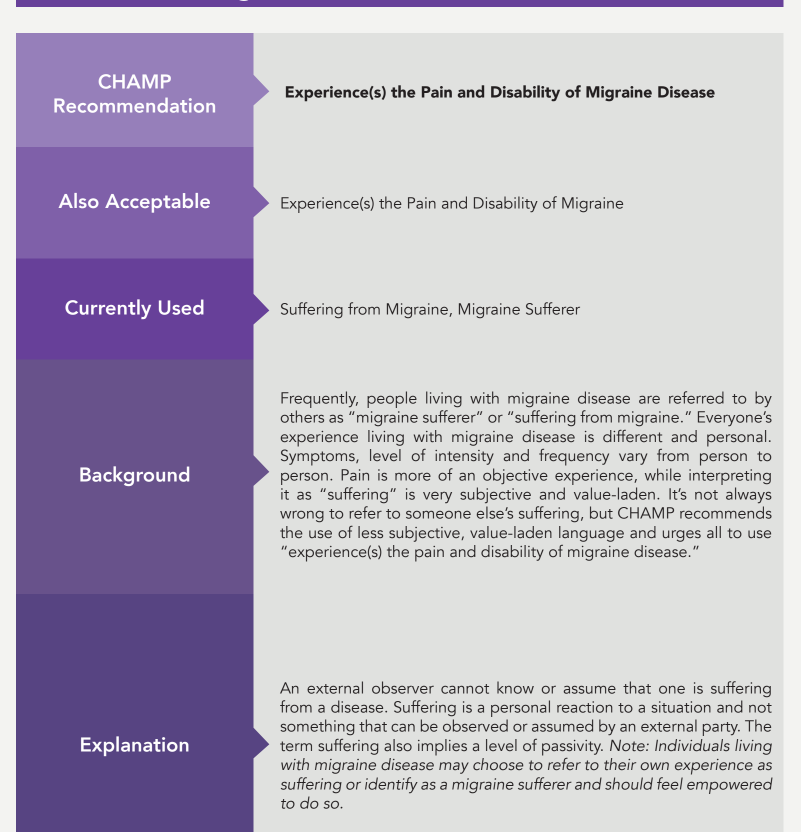
Headache Disease – Overall (continued)



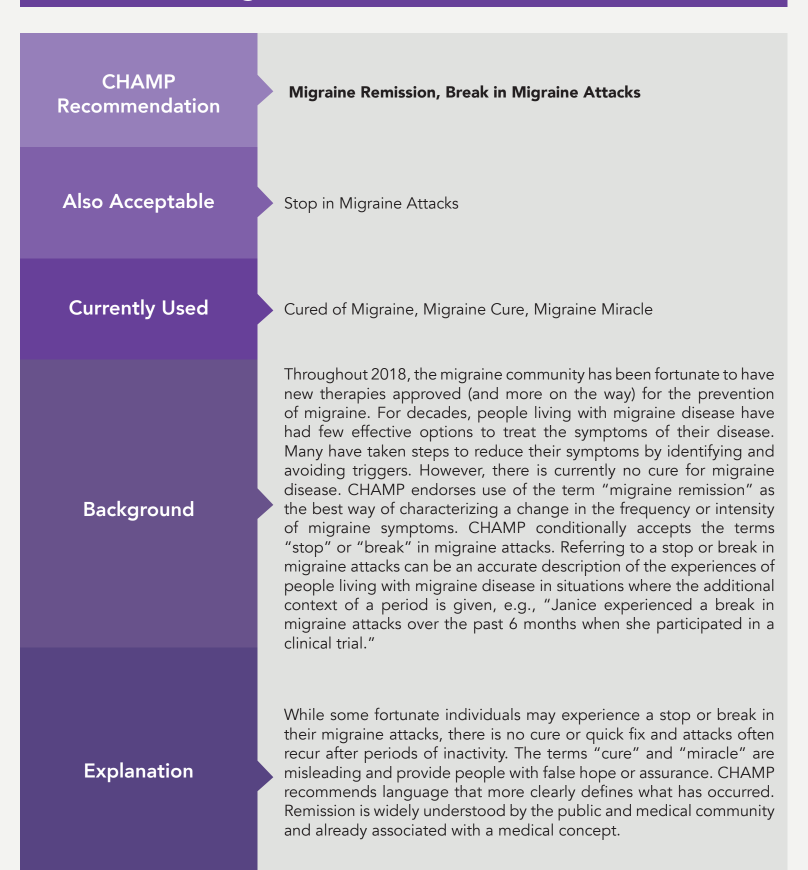
Migraine Disease











Migraine Disease – For Further Discussion





Cluster Disease

Cluster disease, a form of Trigeminal Autonomic Cephalgia (TAC), is less common than migraine disease, but more prevalent than most people realize. An estimated 1 in 1,000 Americans experience cluster attacks. Cluster attacks are frequently described as the most painful physical experience a person can have. Cluster disease is sometimes referred to as "suicide headaches" because, sadly, the suicide rate for this patient population is 20 times higher than the national average. This heightened suicide rate among people with cluster disease is largely believed to occur because of the intensity of the pain they have been living with for extended periods of time.

The cluster community has shared a diversity of opinions on the terminology they feel most appropriately should be used to describe their disease. This is understandable since for decades medical experts have struggled to find common ways of describing or diagnosing cluster disease. Consequently, CHAMP does not currently make language recommendations regarding cluster disease. Instead, CHAMP is providing a listing of the most commonly used cluster terms and the pros/cons of using these terms.

Cluster

PROS

The term "cluster" is simple yet descriptive of the nature of the disease, with attacks often happening in bunches.

CONS

Cluster does not depict the severity of the disease and lacks any medical association. It also overlaps with many other common uses of the word such as to describe a group of people.

Cluster Disease (continued)

Cluster Headache

PROS

"Cluster headache" is more detailed and narrowly focused on the disease. This term is what is currently used under the IHS classification and most prevalent in online communications.

CONS

Symptoms of attacks include more than just headache pain and the severity of the pain is difficult for other people to understand when their only reference point is their own experience of more common headaches.

Cluster Disease

PROS

Using "cluster disease" helps to reinforce this as a serious, medical condition

CONS

Because cluster disease is not yet commonly used, it could be challenging to change the language and build awareness both inside and outside of the community.

TAC (Trigeminal Autonomic Cephalgia)

PROS

Trigeminal Autonomic Cephalgia is a technical, medical and serious depiction of the disease.

CONS

The term can be difficult to say and remember, and it is not widely used. It is also not entirely accurate because cluster is a prevalent type of condition under the broader TAC category, which also includes other conditions.

Suicide Headache

PROS

According to statistics, "suicide headache" is a numerically-justifiable name for the disease. It is attention grabbing and reinforces the critical nature of this condition.

CONS

The use of this term can be perceived as overly sensational and carries a risk of creating despair amongst those who have this condition and possibly contributing to increased suicide rates.



Image Guide

Just as the words we use greatly impact perceptions, so do the images we choose. CHAMP asks all communicators to be thoughtful when selecting imagery and photographs of people with headache diseases to use in publications, advertisements, blogs and websites. The images selected should accurately and honestly depict what it's like to live with these diseases and the range of how they impact people's daily lives.

Images commonly used in the media most frequently portray only white, middle-aged women showcasing mild head pain. The reality is headache, migraine and cluster diseases do not discriminate. They impact all ages, races and genders. And the symptoms during an attack far exceed just head pain. Most people also experience other symptoms including nausea, vomiting, cognitive impairment, visual disturbances and more.

On the following pages, we have included examples of photos that are commonly used to illustrate the limited representation that is prevalent in current communications. We have also included examples of photos that provide more accurate and diverse depictions of headache, migraine and cluster disease. CHAMP advocates for the use of images similar to these more diverse and accurately representative examples.



CURRENTLY COMMON IMAGES – DO NOT USE EXCLUSIVELY







Head pain is NOT the only symptom



We are NOT weak

The following images show how headache and migraine are commonly portrayed using white, middle-aged, women. If this demographic is shown, it should be paired with other images that show more diverse demographics impacted by headache diseases.



SUGGESTED IMAGE EXAMPLES





Exhaustion and sensitivity to light

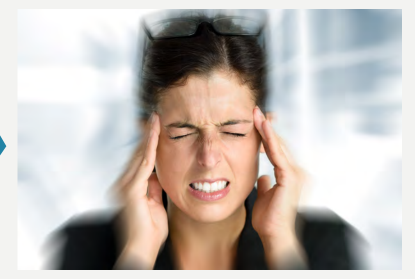
CHAMP endorses the following images as examples of preferred images because they show the wider array of associated symptoms and varied ages, gender, races and ethnicities impacted by headache diseases.



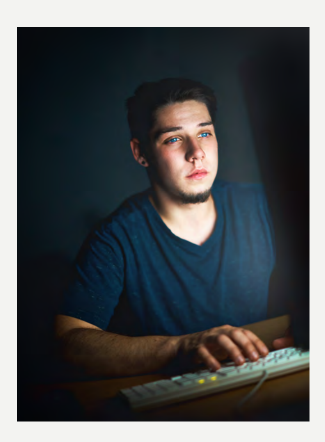
Depression and anxiety







SUGGESTED IMAGE EXAMPLES







Migraine often decreases with age, but not always CHAMP endorses the following images as examples of preferred images because they show the wider array of associated symptoms and varied ages, gender, races and ethnicities impacted by headache diseases.



Headache diseases impact men



SUGGESTED IMAGE EXAMPLES



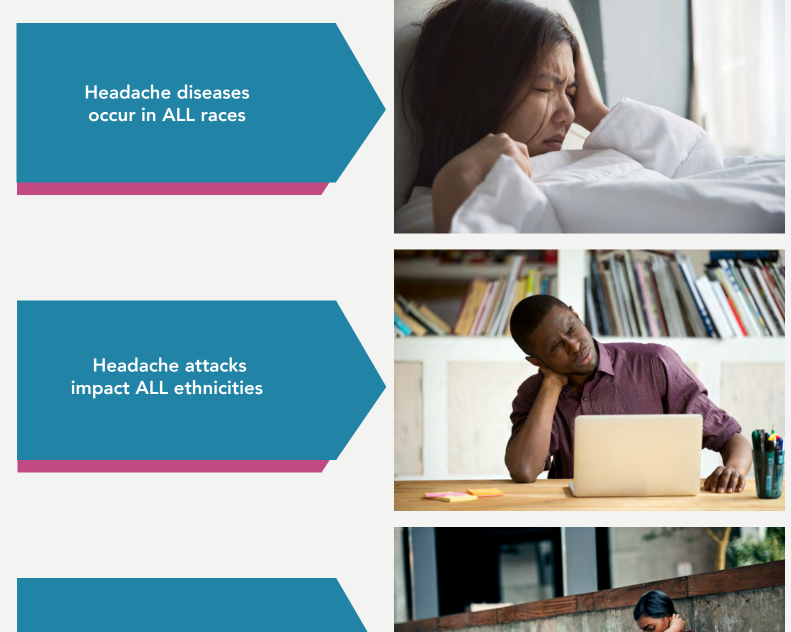
Headache attacks also interrupt the work of men



Headache attacks lead to noise sensitivity in men also



Headache attacks lead to light sensitivity in men also CHAMP endorses the following images as examples of preferred images because they show the wider array of associated symptoms and varied ages, gender, races and ethnicities impacted by headache diseases.



Headache diseases do not discriminate



SUGGESTED IMAGE EXAMPLES



Headache diseases are disproportionately prevalent in members and veterans of the military



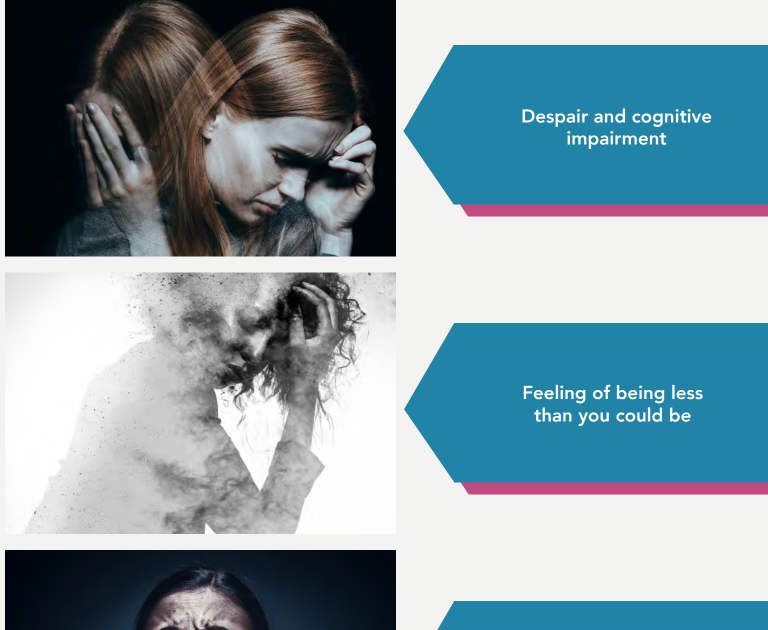
Headache diseases are disproportionately prevalent in members and veterans of the military

POWERFUL IMAGES

People are bombarded by communications, messages and visuals from mass media and social media. Those working to share educational messages about chronic disease are competing against major advertisers and entertainment providers that use images and messages that are captivating and visually compelling. CHAMP believes there are ways to educate all stakeholders on headache, migraine and cluster disease that use engaging, accurate, inclusive and empathetic visuals. The following images are examples of the compelling visuals we recommend.



SUGGESTED POWERFUL IMAGE EXAMPLES



Pair



Conclusion

This document represents consensus guidance from CHAMP member organizations and the headache disease patient communities we serve. We reiterate that the purpose of this guide is to help all people who communicate about headache diseases to do so in a more accurate, informed and empathetic manner. This guide should only be used as an educational tool and never to shame anyone for the language they use.

Language is constantly evolving and CHAMP will update this guide over time so that it reflects the most recent consensus of the headache patient advocacy community. If you have any feedback or suggestions for this guide, please email Kevin Lenaburg, Executive Director of CHAMP, at Kevin dot Lenaburg @ HeadacheMigraine dot org.

¹ Young, W. B., MD, Kempner, J., PhD, Loder, E.W., MD, Roberts, J. PhD, Segal, J.Z., PhD, Solomon, M., PhD, Cady, R.K., MD, Janoff, L., BA, Sheeler, R.D, MD, Robert, T., PhD, Yocum, J., RN, Sheftell, F. D., MD (2011). Naming Migraine and Those Who Have It. Headache, 0017-8748, 283-291.

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