October 17, 2023

The Honorable Xavier Becerra  
Secretary  
U.S. Department of Health and Human Services  
200 Independence Avenue, SW  
Washington, DC 20201

The Honorable Lisa M. Gomez  
Assistant Secretary  
Employee Benefits Security Administration  
U.S. Department of Labor  
200 Constitution Avenue, NW  
Washington, DC 20002

The Honorable Douglas W. O’Donnell  
Deputy Commissioner for Services and Enforcement  
Internal Revenue Service  
U.S. Department of the Treasury  
1111 Constitution Avenue, NW  
Washington, DC 20224

Re: Comments on Technical Release 2023-01P

Dear Secretary Becerra, Assistant Secretary Gomez, and Deputy Commissioner O’Donnell;

Our organizations, who represent consumers, family members, mental health and substance use disorder (MH/SUD) providers, advocates, and other stakeholders, appreciate the opportunity to comment on the Department of Health and Human Services, Employee Benefits Security Administration, and the Internal Revenue Service’s (the “Departments”) Technical Release 2023-01P, Request for Comment on Proposed Relevant Data Requirements for Nonquantitive Treatment Limitations (NQTLs) Related to Network Composition and Enforcement Safe Harbor for Group Health Plans and Health Insurance Issuers Subject to the Mental Health Parity and Addiction Equity Act (hereinafter "Technical Release").

We strongly support the Departments’ proposed NQTL data collection requirements relating to network composition as part of the Departments’ efforts to increase access to MH/SUD treatment. Such data collection is critical to ensure that plans and issuers do not impose treatment limitations that place a greater burden on plan members’ access to MH/SUD treatment than to medical/surgical (M/S) treatment. Combined with the accompanying proposed requirements related to the Mental Health Parity and Addiction Equity Act (MHPAEA), the data collection requirements that are envisioned in the Technical Release would be powerful steps in the right direction to increasing access to MH/SUD treatment. We urge the Departments to require that the data points for MH
services and SUD services be separately collected, analyzed and reported, consistent with MHPAEA statutory and regulatory requirements. Data should also be collected for M/S services to facilitate MHPAEA comparisons. We also urge the Departments to require that all data be collected, analyzed, and reported by age group, including children and adolescents, and by race/ethnicity (where possible). The Departments should also develop uniform definitions and methodologies for the collection of all data points so that valid data are collected and can be compared across plans/issuers.

We appreciate the Departments’ commitment to ensuring that the data plans/issuers will be required to collect are an accurate reflection of individuals’ access to treatment. Given that the Departments’ guidance to plans will likely need to evolve over time to ensure such accuracy, we urge the Departments not to proceed with a “safe harbor” for plans/issuers based on data collection that has yet to be validated as meaningful. As we describe below, we believe that a “safe harbor” should not be explored until data collection has been extensively validated. Otherwise, the Departments may give “safe harbor” to plans/issuers that impose discriminatory barriers that inhibit access to MH/SUD treatment.

Our full comments are as follows.

**Out-of-Network Utilization**

Studies indicate that the percentage of services received out of network (OON) is a key indicator of the availability of in-network services. Due to the higher cost-sharing of OON services, individuals rarely choose to obtain care OON if adequate in-network services are available on a timely basis. The landmark Milliman report demonstrates the importance of such data and how frequently MH/SUD care is obtained OON compared to M/S care. The data should be disaggregated by age groups wherever possible, so that utilization by children and adolescents can be distinguished from adults. This is particularly important given that half of lifetime mental health conditions begin by age 14 and our country’s ongoing youth mental health emergency.

We support the Departments’ reference to quantitative templates in the Appendix that have already been validated and are in use by employer groups and state regulators. The Bowman Family Foundation Report, which is based on a patient and provider survey conducted by NORC, shows multiple analyses of OON use and access problems, as do other consumer and employer and provider surveys and studies. Recently published research also shows that MH/SUD patients go out of network because of MH/SUD network inadequacies. This research also found that M/S patients go out of network for the similar reasons, yet MH/SUD patients go out of network at a much higher rate due to the significantly higher rate of MH/SUD network inadequacies compared to M/S.
**Percentage of In-Network Providers Actively Submitting Claims**

Research indicates that collecting this data is critically important to determining the adequacy of a network. Plans/issuers frequently pad their networks by having providers listed as in-network even if they aren’t actively submitting claims. This metric can also be important in suggesting the existence of other reasons why providers listed as in-network might not be available, including low reimbursement that incentivizes providers to fill appointments with patients with insurance that pays more and/or cash-pay patients. Again, this data should be disaggregated by children and adolescents wherever possible. While we welcome the Departments’ reference to child psychiatrists and psychologists, all types of pediatric providers should be included. Additionally, it is important to include data on M/S pediatric subspecialists to the lists (e.g., pediatric cardiologists, pediatric neurologists, etc.) for purposes of assessing parity. We encourage the Departments to require actual participation data on all sub-types of MH/SUD professional providers for both adults and children, as well as inpatient and outpatient facilities.

**Time and Distance Standards**

We strongly support the Departments’ suggestion that the Departments collect detailed data on the percentage of participants/beneficiaries/enrollees who can access specified provider types in-network within a certain time and distance. We strongly agree with the Departments’ view that this data would help with the assessment of a plan/issuer’s operational compliance with respect to any NQTLs related to network composition. We also recommend that the Departments collect data on appointment wait times, which are an essential metric to measure network adequacy and the most critical for participants/beneficiaries seeking timely access to care. The Department of Health and Human Services has already put forward strong proposed standards for Medicaid managed care and the Children’s Health Insurance Program (CMS-2439-P), which establish maximum appointment wait time standards for routine outpatient mental health and substance use disorder services of 10 business days and require such independent secret shopper surveys. These standards align with appointment wait time metrics that have been adopted for Qualified Health Plans. We recommend that any network adequacy standards, such as time/distance, wait times, etc., issued by state or federal governments identify key sub-types of MH/SUD professional providers, such as child and adult psychiatrists, child and adult psychologists, master’s level social workers and mental health counselors, psychiatric ARNPs, psychiatric PAs. In addition, all acute and sub-acute inpatient sub-types should have specific network adequacy standards, as well as sub-types of outpatient facility programs, such as IOP, PHP, ABA, MAT, eating disorder, etc.

In collecting data, the Departments should collect data on routine and crisis appointments, including for follow-up and ongoing care. When only initial appointment wait times are measured, plans/issuers can manipulate their practices to have initial “intake” appointments while having long delays in the delivery of ongoing services. Data
should be disaggregated by age group to assess wait times and travel distance for children and adolescents.

We also urge the Departments to require any plan/issuer that uses a source or evidentiary standard for its network adequacy standards (whether a state/federal government or an independent entity such as NCQA) to identify and explain how the standards were designed, as written, to comply with MHPAEA. The Departments should require that, for any source, a plan/issuer must provide and define all the factors and evidentiary standards relied upon for each MH/SUD network standard (e.g., time and distance) and complete a comparative analysis for each factor to demonstrate that the standard is comparable and no more stringent, as designed, for MH/SUD than for M/S.

For example, MH/SUD outpatient providers often have different characteristics such as smaller size and/or smaller caseloads than M/S providers. It is essential that the Departments require plans/issuers to demonstrate that these different characteristics are considered and addressed in assessing the adequacy of each standard. As an illustration, many MH/SUD professionals can only treat 8 to 10 patients per day, while many Primary Care Physicians (PCPs) can see 30 to 40 patients per day. A network adequacy standard that has equivalent time and distance standards (10 miles / 30 minutes) for one full-time PCP and one full-time Psychologist is not comparable and is more stringent, due to the provider case load.

The Departments should require the same type of analysis for MH/SUD facilities. For example, how are MH/SUD acute and subacute inpatient facilities the same or different as compared to acute and subacute M/S facilities – and how is that considered and addressed by the plan in developing each standard? The plan should be required to describe the factors used to compare types of MH/SUD facilities (e.g., psychiatric versus substance use), as well as capacity (e.g., number of beds, availability of beds) of MH/SUD facilities versus M/S facilities.

We urge the Departments to also ensure that as-written NQTL analysis also address the factors of supply/demand for both MH/SUD and M/S outpatient professional and facility providers, including definitions for these factors, evidentiary standards and sources. Studies, reports or data measuring provider supply (including shortages) and market demand should be required to be provided.

Network Availability and Distribution of Professions

We applaud the Departments for focusing on whether providers are accepting new patients (Section (c)(4)(iv)(A)(2) of the proposed rule), which is a crucial issue in light of the high demand for MH/SUD services. Given the high demand for MH/SUD services, very few of these providers will have wide open availability to take new patients. Thus, we recommend including a “limited availability” category to provide clearer information.
on availability.\textsuperscript{1} A MH/SUD provider with just a few time slots available does not add significant capacity to plans/issuers’ networks. We believe that the Departments should require that any network adequacy standard should consider typical limits on MH/SUD providers, who typically have smaller caseloads, less capacity and limited availability for new patients as compared to most M/S professional providers. (For example, a standard that equates 1 full-time PCP to 1 full-time Psychologist is not comparable in light of the differences in caseloads and capacity).

It is also important to require metrics on the number of available providers who fill high-demand needs in the network, such as those seeing children & adolescents, those who specialize in eating disorders or LGBTQ patients, and those who meet the language needs of the population served by the network. While the Service Utilization metrics below in these same categories would address how much certain services are being utilized, it may be that while there is a reasonable level of, for example, eating disorder services provided by network providers, those providers may be completely full. Thus, it is also important to assess whether new patients with these specialized needs can find available providers.

A robust network has a full range of different professions and training levels to handle the varying needs and more complex problems of the patient population. Thus, we recommend gathering data (on both the MH/SUD and M/S sides) on the percentage of the top 10 different professions that make up the network. We also support that plans should measure the actual numbers of licensed MH/SUD professionals by geo zip code.

**Network Admissions**

In assessing network composition and access to MH/SUD services, we urge the Departments to review the criteria and processes by which plans/issuers determine which providers to admit into networks and/or how plans/issuers define when a network is considered “full” or “closed.” Reports from MH/SUD providers suggest that they are often denied participation on networks due to the networks being “closed” or “full,” even though patients are unable to find appropriate providers in that network. Other providers who are eventually admitted into networks report having to wait as long as nine months to be added.

Plans/issuers should not be allowed to claim a workforce shortage as a reason for access to care issues and simultaneously keep networks locked or slow to accept new providers. Measuring and monitoring access to care for all sub-types of MH/SUD providers will reveal how much responsibility plans/issuers bear for the lack of access to MH/SUD services. For example, plans/issuers should provide metrics on how many providers applied to the network, what percentage were rejected and the reasons for the rejection (e.g., network full, provider not qualified, and the time it takes to bring providers into the network from when they first apply).

\textsuperscript{1} This concept comes from the following: Massachusetts Division of Insurance, Report of the Provider Directory Task Force to respond to Section 4 of Chapter 124 of the Acts of 2019, (2020), available at: https://www.mass.gov/doc/provider-directory-task-force-report-2020/download
Reimbursement Rates

We applaud the Departments’ suggested data collection relating to reimbursement rates, which are critical determinants of network adequacy; many studies show the strong correlation between network access and reimbursement rates. We also commend the Departments for putting forward potential requirements that reimbursement rate data be “compared to billed rates.” Reimbursement rates that are not reflective of current market reimbursement can profoundly affect the availability of MH/SUD providers, including current providers’ decision to join a network and potential providers’ decisions whether to enter the field. We strongly recommend the Departments evaluate the ratio of allowed in-network and OON amounts to OON billed market rates for MH/SUD and M/S. The billed rates of OON providers are the most accurate representation of the market rate. We also support developing additional reimbursement rate measures, such as percent of out-of-pocket (OOP) expenses for enrollees using out-of-network providers for MH/SUD versus M/S care.

With respect to the use of Medicare Fee Schedule and other external benchmarks such as Fair Health, we urge the Departments to utilize significant care to avoid perpetuating historic (and ongoing) disparities between MH/SUD and M/S reimbursement rates that are embedded in these benchmarks. We urge the Departments to recognize that Medicare and other claims databases and benchmarks rely on historical data that embeds legacy disparities in reimbursements between MH/SUD and M/S. Additionally, we strongly believe that caution is warranted with respect to Medicare because it:

- Is not subject to MHPAEA;
- Does not have allowed amounts for certain sub-types of MH/SUD providers (e.g., sub-acute inpatient care and the full range of MH/SUD professional providers and psychosocial rehabilitation services);
- Does not cover some MH/SUD services for children and adolescents given that this population does not participate in the program; and
- Has a structure that undervalues the work of MH/SUD professionals, which CMS recently acknowledged in its recent Physician Fee Schedule proposed rules.

Nonetheless, we recognize that the Departments, multiple state regulators, and research organizations (such as Milliman) have documented significant disparities between Medicare allowed amounts and plans/issuers’ allowed amounts for MH/SUD providers versus M/S providers. As described below, the ultimate determiner of parity for any reimbursement comparison is the access to services (i.e., adequacy) within MH/SUD networks in comparison with M/S networks. Indeed, reimbursement rate comparisons could actually show that MH/SUD providers are reimbursed at the same level as M/S providers, yet if MH/SUD network inadequacies persist, plans/issuers should be required to increase rates further for MH/SUD providers to address network inadequacies, as plans/issuers do for M/S.

While taking into account that the Medicare fee schedule and other external benchmarks may have legacy disparities embedded for MH/SUD services compared to
M/S services, we have seen that they can be used as tools to demonstrate parity non-compliant reimbursement rates. This was the case in the U.S. Department of Labor and New York Attorney General’s 2021 lawsuit against United Healthcare and United Behavioral Health (UBH) and resulting settlement agreement, which were based, in part, on UBH’s disparate reductions from baseline rates derived from Medicare. The Departments have made it clear that when faced with M/S provider shortages, if plans increase reimbursement rates for M/S providers to ensure adequate M/S networks, they must increase rates to address MH/SUD providers shortages as well to ensure adequate behavioral networks. The Bowman Family Foundation publication, “Federal Parity Law (MHPAEA): NQTL of In-Network Reimbursement Rates: Non-Comparable Use of Factors of Provider Leverage a/k/a Bargaining Power and Workforce Shortages” references federal data that shows there are more zip codes in the U.S. with PCP shortages than Psychiatrist shortages. Yet, there is relatively low out-of-network use for PCPs, and PCPs are routinely paid more than Psychiatrists for the same evaluation and management billing codes. Key quotes include:

“Well, on average, the average in-network reimbursement for MH/SUD professional office visits from commercial insurers was approximately 2.5% below Medicare reimbursement, and OON use of such visits was approximately 17%, i.e., 5.4 times higher than for primary care providers.”

“Well, on average, the average in-network reimbursement for primary care professional office visits from commercial insurers was approximately 20% above Medicare reimbursement, and OON use of such visits was approximately 3%.”

“HRSA identifies “Health Provider Shortage Area” (HPSA) designations, which indicate that demand far exceeds supply. As reported by Kaiser Family Foundation, this national data as of Sept. 30, 2021 shows more shortages for PCPs than for mental health providers (7447 vs. 5930 shortage areas).”

The Department’s guidance in the 2020 Self Compliance Tool is also clear:

“NOTE – Plans and issuers may attempt to address shortages in medical/surgical specialist providers and ensure reasonable patient wait times for appointments by adjusting provider admission standards, through increasing reimbursement rates, and by developing a process for accelerating enrollment in their networks to improve network adequacy. To comply with MHPAEA, plans and issuers must take measures that are comparable to and no more stringent than those applied to medical/surgical providers to help ensure an adequate network of MH/SUD providers, even if ultimately there are disparate numbers of MH/SUD and medical/surgical providers in the plan’s network…” (Emphasis added).

As with all quantitative data metrics, multiple measures are important to accurately assess the compliance of any NQTL. Consistent with the current regulations and enforcement, as well as the Proposed Rules, reimbursement rates for MH/SUD providers are a key aspect of in-network access to care. We have seen that plans/issuers use reimbursement rate increases to establish and maintain adequate
M/S networks, especially in addressing shortages of M/S providers. MHPAEA requires plans to take the same measures for MH/SUD providers to ensure adequate networks.

Aggregate Data Collection

We strongly support the Departments, when reviewing self-funded employer group plans, to require relevant data to be collected and evaluated for both employer group enrollees as well as enrollees of the employer’s third-party administrator (TPA) or other service provider in the aggregate. We agree with the Department that individual employer group plans may lack sufficient data.

Service Utilization Data

In assessing network composition and access to MH/SUD services, we urge the Departments to require plans to report on utilization rates for specific MH/SUD services and level of care. These utilization rates should be compared to estimates of participants/beneficiaries with these conditions, as well as utilization rates for M/S services. Examples of services providers, settings, and levels of care on which we urge the Departments to collect utilization data include:

- Child and adult psychiatrists, child and adult psychologists, master’s level social workers and mental health counselors, psychiatric ARNPs, psychiatric PAs, all acute and sub-acute inpatient sub-types, and sub-types of outpatient facility programs, such as IOP, PHP, ABA, MAT, eating disorders, etc.;
- Each of the levels (and sub-levels) of care described in The American Society of Addiction Medicine (ASAM) Criteria and the age-specific Level of Care Utilization System (LOCUS) family of criteria developed by the American Association of Community Psychiatrists and the Academy of Child and Adolescent Psychiatry, as well as the average length of stay / treatment units and denial rates by each of these levels of care;
- Service utilization by MH/SUD diagnoses;
- High-demand needs such as services for children and adolescents, eating disorder, and services by providers who meet the language needs of the population served by the network;
- Cognitive behavioral therapy;
- Dialectical behavioral therapy;
- Coordinated Specialty Care;
- Medications for opioid use disorder (MOUD);
- Medications for alcohol use disorder (MAUD); and
- Medications for bipolar disorder, schizophrenia, major depressive disorder, and other MH/SUDs.

While there are numerous underutilized MH/SUD services, particularly glaring examples include:
• **Coordinated Specialty Care for First Episode Psychosis.** More than 110,000 people each year experience symptoms of psychosis for the first time (first episode psychosis, FEP) each year. Yet Coordinated Specialty Care (CSC), the evidence-based intervention for FEP that has been heavily researched by the National Institute of Mental Health’s Recovery After Initial Schizophrenia Episode (RAISE) Initiative, is rarely covered by commercial insurance. The result is an underdeveloped treatment system that relies almost exclusively on taxpayer funding. Tragically, estimates suggest that fewer than one-quarter (and likely only about one-tenth) of individuals experiencing FEP receive this life-changing services. Plans/issuers have simply been allowed to offload their responsibility to cover CSC services for FEP to taxpayers in a way that would simply be unimaginable for a physical health condition like Type I diabetes, which similarly often first presents among youth. The Departments must take action to measure and advance access to CSC for individuals experiencing FEP, particularly given that CMS just took a very important step of creating two HCPCS codes for CSC (H2040 which can be billed per month and H2041 which can be billed per encounter).

• **Treatment (ACT).** ACT is a team-based service delivery model designed to help adults with serious mental illness, designed to help individuals recover and live in the community. According to SAMHSA, it is “one of the oldest and most researched evidence-based practices for treatment people with serious mental illness.” It is intended to assist individuals who experience frequent hospitalizations, psychiatric crises, substance use, involvement in the criminal justice system, or homelessness. Yet this service is incredibly underutilized, and almost never available via commercial plans, forcing individuals who could otherwise be supported in the community into more restrictive settings. While data is available on provision of ACT services by state mental health authorities, no such comparable data is available from plans. The Departments should require plans/issuers to collect and analyze ACT utilization data.

• **MH/SUD Emergency Services for MH/SUD Crises.** Federal policymakers have dedicated enormous effort to standing up the 988 Suicide and Crisis Lifeline and expanding MH/SUD crisis services, which help people get the help they need and avoid needless, and often tragic, encounters with law enforcement. To our knowledge, essentially all plans/issuers cover EMS and emergency transport services (which are included in all state benchmark plans), but extraordinarily few plans/issuers cover mobile crisis response team services for MH/SUD emergencies. This failure means that many individuals do not have appropriate coverage of these services. While we encourage the Departments to make clear in the proposed MHPAEA rules that MHPAEA requires all MH/SUD emergency services to be covered when physical health emergency services are covered, it is also critical that the Departments require plans to collect and analyze utilization data on MH/SUD emergency services, including mobile crisis response teams and crisis receiving and stabilization.

• **Dialectical Behavioral Therapy for Various Conditions.** Dialectical Behavior Therapy (DBT) is one of the most effective treatments for self-harm and suicidality and can be useful to treat borderline personality disorder and other
mental health condition. Yet its utilization is very low. The Departments should require plans/issuers to collect and analyze DBT utilization data.

- **Medications for Opioid Use Disorder (MOUD).** Tragically, annual overdose deaths – driven in large part by synthetic opioids – increased dramatically during the pandemic and remain at all-time highs of over 110,000. Yet, despite this enormous increase, only about 1 in 5 individuals with opioid use disorder (OUD) receive medications to treat it. While many contributing factors, inadequate health insurance coverage, including the inability of individuals to find in-network providers, is a critical issue. We urge the Departments to collect and analyze data specifically on utilization of MOUD.

- **Medications for Alcohol Use Disorder (MAUD).** The terrible toll of alcohol use disorders (AUD) frequently goes unrecognized. Yet more than 140,000 Americans died each year between 2015-19 from excessive alcohol use. And with increased alcohol use during the pandemic, estimated deaths increased 25% in 2020. Tragically, effective medications to treat AUD are underutilized, with fewer than 1 in 10 individuals with AUD using these medications. To help increase utilization, we urge the Departments to collect and analyze data specifically on utilization of MAUD.

Our organizations acknowledge that there are many factors contributing to this underutilization, but insurance barriers (including oftentimes the complete lack of coverage) are a major contributing factor that must be addressed. The enormous costs of underutilization of MH/SUD services are currently being shifted onto individuals, their families, public programs (including Medicaid), and taxpayers. Such cost shifts should not be accepted, particularly when such a situation would not be tolerated for physical health conditions.

**Safe Harbor**

The Technical Release also requested feedback on the potential of a “safe harbor” for NQTLs related to network composition. We urge the Department not to proceed with a safe harbor at this time. We understand the desire to most effectively target the Departments’ enforcement resources. However, network adequacy has always been difficult to define and easy to mismeasure. Even when plans have been provided with templates by various state regulators, data is often incomplete, inconsistent and/or contradictory. Thus, a safe harbor has the potential to be harmful if the data collection requirements are not capturing a full and complete picture of participants/beneficiaries’ access to MH/SUD services.

Given the significant work that the Departments need to do – and likely refinements that are necessary over time – to ensure collected data is complete, accurate, and meaningful, a safe harbor should not be considered in the near future. Such a safe harbor should only be considered when the Departments and key consumer stakeholders are confident that the data collected accurately captures actual access to MH/SUD services. Data templates should be validated for operational feasibility and accuracy. If a safe harbor is put in place prior to this occurring, it could cause enormous
damage by giving noncompliant plans/issuers a “safe harbor” against accountability. Furthermore, an issuer residing within such a “safe harbor” would almost certainly escape meaningful oversight from any applicable State authority.

**Meaningful Data & Preventing Data Manipulation**

To ensure that the proposed requirements relating to outcomes data and actions to address material differences in access are meaningful, we urge the Departments to issue standardized definitions on all data points and on methods for gathering and reporting data. For example, the Departments propose collecting data on the number and percentage of claims denials. Yet, there are many ways that plans can collect, and potentially manipulate, such “claims denials” data. For example, the Departments should make clear that failure to pay a claim in part or in full constitutes a denial and must find ways to capture common practices of undocumented denials that occur verbally through peer-to-peer reviews. Additionally, plans can manipulate denial data by approving each visit or day of treatment (thereby increasing the denominator) while telling the provider verbally that further visits/days will not be approved, which is another common occurrence. Such practices can result in meaningless data that bears little resemblance to what individual patients experience. The Appendix to the Technical Release lists templates already in use, including the Bowman Family Foundation’s [Model Data Request Form](#), which includes a section on Denial Rates. We support the continued use of templates that address the issues set forth above.

**Disaggregating MH and SUD Data**

We also encourage the Departments to make clear that MH and SUD data must be collected and analyzed separately. When MH and SUD data is simply aggregated, it can hide important discriminatory impacts. MHPAEA’s requirements apply to MH and SUD benefits individually, which must be reflected in its data collection and analysis requirements.

**Conclusion**

We have included numerous citations to supporting research, including direct links to the research. We direct the Departments to each of the materials we have cited and made available through active links, and we request that the full text of each of the studies and articles cited, along with the full text of our comment, be considered part of the formal administrative record for purposes of the Administrative Procedure Act. If the Departments are not planning to consider these materials part of the record as we have requested here, we ask that you notify us and provide us an opportunity to submit copies of the studies and articles into the record.

Thank you for the opportunity to comment on this important issue. If you have further questions, please contact David Lloyd ([david@thekennedyforum.org](mailto:david@thekennedyforum.org)) or Lauren Finke ([lauren@thekennedyforum.org](mailto:lauren@thekennedyforum.org)) at The Kennedy Forum.
Sincerely,

The Kennedy Forum
American Foundation for Suicide Prevention
American Psychological Association Services
American Society of Addiction Medicine
Eating Disorders Coalition
Inseparable
Mental Health America
NAMI - National Alliance on Mental Illness
National Association for Behavioral Healthcare
National Council for Mental Wellbeing
National Health Law Program
Partnership to End Addiction
Policy Center for Maternal Mental Health
Psychotherapy Action Network
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