Catabasis Pharmaceuticals Presents New Edasalonexent Data Showing Significantly Slowed Duchenne Muscular Dystrophy Disease Progression as Measured by MRI Through One Year of Treatment

-- Statistically Significant Improvements in MRI T2 Rate of Change Compared to Control Consistent with Improvements Demonstrated in Assessments of Muscle Function in MoveDMD® Trial --

CAMBRIDGE, Mass., April 25, 2018 – Catabasis Pharmaceuticals, Inc. (NASDAQ:CATB), a clinical-stage biopharmaceutical company, today reported new positive magnetic resonance efficacy results showing slowed disease progression in boys with Duchenne muscular dystrophy (DMD) in the MoveDMD trial through 48 weeks of edasalonexent treatment. Both magnetic resonance imaging (MRI) and magnetic resonance spectroscopy (MRS) were performed in the Phase 2 MoveDMD trial and open-label extension. Statistically significant improvement in the rate of change in lower leg composite MRI T2 at 12, 24, 36 and 48 weeks on oral 100 mg/kg of edasalonexent treatment were observed compared to the off-treatment control period (p<0.05 for all time points). Improvements in changes in both soleus and vastus lateralis (VL) MRS fat fraction through 48 weeks of edasalonexent treatment compared to the off-treatment control period were also demonstrated. These data were presented today at the American Academy of Neurology 70th Annual meeting in Los Angeles, CA. These improvements in MRI T2 and MRS fat fraction show a slowing of disease progression and are in addition to the improvements observed in all assessments of muscle function through more than a year of edasalonexent treatment.

“The results reported to date from the MoveDMD trial have been consistent and support edasalonexent as a disease-modifying therapy. Overall, in the MoveDMD trial edasalonexent has slowed progression of this disease based on improvements in multiple assessments of physical function.”
function and biomarkers of muscle health and inflammation,” said Jill C. Milne, Ph.D., Chief Executive Officer of Catabasis. “We believe that these effects ultimately will translate to boys with Duchenne maintaining functional abilities longer.”

“I am very encouraged to see these emerging data with favorable MRI changes and results of muscle function assessments with edasalonexent, which are clearly divergent from the untreated disease progression that I see in boys in this age range with Duchenne,” said Richard Finkel, M.D., Chief, Division of Neurology, Department of Pediatrics at Nemours Children’s Hospital, Orlando and a Principal Investigator for the study. “There is a strong need for a therapy with a safety and tolerability profile like edasalonexent for the many boys affected by this devastating disease.”

MRI is a non-invasive approach to assess disease progression in DMD. MRI T2 as well as MRS fat fraction and T2 were performed in the Phase 2 MoveDMD trial and open-label extension. MRI T2 measures combined inflammation and fat in one measurement. MRS T2 measures the inflammatory component independently and MRS fat fraction measures the amount of fat in the muscle. As boys with DMD get older, the amount of fat in their muscles increases with consequent loss of functional abilities. Changes in MRI T2 and MRS fat fraction are known to correlate with changes in function in boys with DMD. Increases in both measures strongly correlate with worse performance on timed function tests and predict future loss of functional abilities. FDA considers MRI to be an important supportive early endpoint demonstrating therapeutic effect as mentioned in the recently released FDA guidance on development in DMD.

Statistically significant improvement in the rate of change in lower leg composite MRI T2 at 12, 24, 36 and 48 weeks of oral 100 mg/kg of edasalonexent treatment were observed compared to the off-treatment control period (p<0.05 at all time points). After 48 weeks of 100 mg/kg of edasalonexent treatment, MRS fat fraction of the soleus muscle increased by an average of 0.85%, while in the same boys during the off-treatment control period the average annualized increase was 2.6% per year. Boys from the ImagingDMD natural history study who were largely on steroids had an increase in soleus fat fraction of approximately 3% per year. After 48 weeks of edasalonexent treatment, MRS fat fraction of the VL (one of the quadriceps) increased by 5.9%, while in the same boys during the off-treatment control period the average annualized increase was 10.4% per year. Boys from the ImagingDMD natural history comparison study who were largely on chronic steroids had an increase in VL fat fraction of approximately 7% per year. MRS T2 of soleus and VL decreased from baseline. The changes in the rate of increase in fat fraction in the soleus and VL muscles demonstrate greater stability with edasalonexent treatment in the MoveDMD trial with less of an increase in fat in their muscles compared to the same boys prior to treatment. The increase in fat fraction observed on edasalonexent treatment was also less than that observed in natural history in a different population of boys with DMD.

Consistent with these magnetic resonance data, in the Phase 2 MoveDMD trial and open-label extension, a preservation of muscle function and slowing of DMD disease progression was seen in boys treated with edasalonexent compared to the rates of change during the control period prior to receiving edasalonexent. Through a year of treatment, the 100 mg/kg/day treatment group showed consistent and clinically meaningful improvements in rates of decline compared to rates
of change during the control period across all four assessments of muscle function: the three
timed function tests (10-meter walk/run, 4-stair climb and time to stand), as well as the North Star
Ambulatory Assessment (NSAA), an integrated global assessment of muscle function.

Edasalonexent is being developed as a potential foundational disease-modifying therapy for all
patients affected by DMD, regardless of their underlying mutation. No evidence of side effects or
safety issues common with the current DMD standard of care has been observed after more than
37 patient-years of exposure to edasalonexent. Catabasis is preparing for a single global Phase
3 trial to evaluate the efficacy and safety of edasalonexent for registration purposes, dependent
on raising capital.

**About the MoveDMD Trial**
The MoveDMD trial is investigating the safety and efficacy of edasalonexent in steroid-naïve boys
enrolled at ages 4 – 7 affected with DMD (any confirmed mutation). The trial is comprised of three
parts - Phase 1, Phase 2 and open-label extension. Phase 2 was a randomized, double-blind,
placebo-controlled 12-week portion with 31 ambulatory boys across a range of dystrophin
mutations. The 12-week MRI T2 primary endpoint for treated boys compared to placebo was
directionally positive although not statistically significant. The open-label extension evaluated
longer term safety and efficacy using pre-specified analyses comparing the rates of change in
boys receiving edasalonexent treatment and prior to treatment. In the Phase 2 and open-label
extension of the MoveDMD trial, edasalonexent substantially slowed DMD disease progression
in boys on 100 mg/kg/day up to 60 weeks of treatment. Across all assessments of muscle function,
consistent improvements were observed in the rate of decline after 12, 24, 36, 48 and 60 weeks
of oral 100 mg/kg/day edasalonexent treatment compared to the rate of change in the control
period for boys prior to receiving edasalonexent treatment. Statistically significant improvements
were also seen in non-effort based measures of muscle health (muscle enzymes and c-reactive
protein and MRI T2 compared to control). In the 100 mg/kg/day treatment group, 16 boys
commenced edasalonexent either at the beginning of Phase 2 or at the beginning of the open-
label extension. Edasalonexent has been well tolerated with no safety signals observed in the
trial.

**About Edasalonexent (CAT-1004)**
Edasalonexent (CAT-1004) is an investigational oral small molecule that is being developed as a
potential disease-modifying therapy for all patients affected by DMD, regardless of their
underlying mutation. Edasalonexent inhibits NF-kB, a protein that is activated in DMD and drives
inflammation and fibrosis, muscle degeneration and suppresses muscle regeneration.
Edasalonexent continues to be dosed in the open-label extension of the MoveDMD Phase 2
clinical trial and Catabasis is preparing for a single global Phase 3 trial to evaluate the efficacy
and safety of edasalonexent for registration purposes, dependent on raising capital. The FDA has
granted orphan drug, fast track and rare pediatric disease designations and the European
Commission has granted orphan medicinal product designation to edasalonexent for the
treatment of DMD. For a summary of clinical results reported to-date, please visit
About Catabasis
At Catabasis Pharmaceuticals, our mission is to bring hope and life-changing therapies to patients and their families. Our SMART (Safely Metabolized And Rationally Targeted) Linker drug discovery platform enables us to engineer molecules that simultaneously modulate multiple targets in a disease. Our lead program in development is edasalonexent for the treatment of Duchenne muscular dystrophy. For more information on edasalonexent and our pipeline of drug candidates, please visit www.catabasis.com.

Forward Looking Statements
Any statements in this press release about future expectations, plans and prospects for the Company, including statements about future clinical trial plans including, among other things, statements about the Company’s plans to commence a single global Phase 3 trial in DMD to evaluate the efficacy and safety of edasalonexent for registration purposes, and other statements containing the words “believes,” “anticipates,” “plans,” “expects,” “may” and similar expressions, constitute forward-looking statements within the meaning of the Private Securities Litigation Reform Act of 1995. Actual results may differ materially from those indicated by such forward-looking statements as a result of various important factors, including: uncertainties inherent in the initiation and completion of preclinical studies and clinical trials and clinical development of the Company’s product candidates; whether interim results from a clinical trial will be predictive of the final results of the trial or the results of future trials; expectations for regulatory approvals to conduct trials or to market products; the Company’s ability to obtain financing on acceptable terms and in a timely manner to fund the Company’s planned Phase 3 trial of edasalonexent in DMD for registration purposes; availability of funding sufficient for the Company’s foreseeable and unforeseeable operating expenses and capital expenditure requirements; other matters that could affect the availability or commercial potential of the Company’s product candidates; and general economic and market conditions and other factors discussed in the “Risk Factors” section of the Company’s Annual Report on Form 10-K for the year ended December 31, 2017, which is on file with the Securities and Exchange Commission, and in other filings that the Company may make with the Securities and Exchange Commission in the future. In addition, the forward-looking statements included in this press release represent the Company’s views as of the date of this press release. The Company anticipates that subsequent events and developments will cause the Company’s views to change. However, while the Company may elect to update these forward-looking statements at some point in the future, the Company specifically disclaims any obligation to do so. These forward-looking statements should not be relied upon as representing the Company’s views as of any date subsequent to the date of this release.

###

Investor and Media Contact
Andrea Matthews
Catabasis Pharmaceuticals, Inc.
T: (617) 349-1971
amatthews@catabasis.com