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	NCT Number	Title	Other Names	Status	Conditions	Interventions	Characteristics	Population	Sponsor/Collaborat	Dates
1	NCT0555 1494	Urinary Concentration of Phthalate Metabolites in Women With and Without Endometriosis Study Documents:	Title Acronym: Other Ids: 2706	Recruiting	Endometriosis	Diagnostic Test: Urinary sample At study entry, we will collect urinary samples from women. Participants will be asked with an interview on demographic and lifestyle characteristics, health-related behaviours, the existence and duration of infertility, medical history, and history of hormonal or surgical treatments for endometriosis. Pain symptoms will be evaluated through a 10 cm long one-dimensional visual-analogue scale (VAS). In addition, women will be asked to report about their habits about consumption of plastic	Study Type: Observational Phase: Study Design: Observational Model: Case-Control Time Perspective: Prospective Primary Outcome Measures: Same as current Secondary Outcome Measures: Same as current	Actual Enrollment: Estimated Enrollment: 440 Original Estimated Enrollment: Same as current Age: 18 Years to 45 Years (Adult) Sex: Female	Study Sponsors: Same as current Collaborators: Not Provided	Study Start: March 22, 2022 Primary Completion: September 30, 2023 (Final data collection date for primary outcome measure) Study Completion: March 30, 2024 First Posted: September 22, 2022 Results First Posted: Last Update Posted: September 22, 2022
2	NCT0554 6229	Assessment of Methadone and Buprenorphine in Interstitial Fluid Study Documents:	Title Acronym: Other Ids: 1332356 5R44DA044905- 03 (U.S. NIH Grant/Contract)	Recruiting	Opioid Use Disorder	Procedure: Microneedle based interstitial fluid collection Interstitial fluid will be collected from the skin using microneedles and suction.	Study Type: Observational Phase: Study Design: Observational Model: Case-Only Time Perspective: Prospective Primary Outcome Measures: Same as current Secondary Outcome Measures: Same as current	Actual Enrollment: Estimated Enrollment: 22 Original Estimated Enrollment: Same as current Age: 18 Years and older (Adult, Older Adult) Sex: All	Study Sponsors: Same as current Collaborators: National Institutes of Health (NIH) National Institute on Drug Abuse (NIDA)	Study Start: September 14, 2022 Primary Completion: October 15, 2022 (Final data collection date for primary outcome measure) Study Completion: October 30, 2022 First Posted: September 19, 2022 Results First Posted: Last Update Posted: September 19, 2022

2 North Authority The Processing of Control Pr		NCT Number	Fitle	Other Names	Status	Conditions	Interventions	Characteristics	Population	Sponsor/Collabora	t Dates
Specific patterns of gene expression related to each metabotype will be investigated in PBMCs by using a microarray-based approach. Analysis will be carried out in a subset of 10 samples for each metabotype.	3	NCT0541	Aggregate Metabolic Phenotypes for (Poly)Phenols: Development of an Oral (Poly)Phenol Challenge Test (OPCT) Study	Title Acronym: Other Ids: 1352/2020/SPER		Individual Variability in (Poly)Phe nol Metabolis m Cardiomet abolic	Dietary Supplement: Oral (poly)phenol challenge test (OPCT) Nutritional challenge with standardized (poly)phenol-rich	Study Design: Allocation: N/A Intervention Model: Single Group Assignment Masking: None (Open Label) Primary Purpose: Screening Primary Outcome Measures: • Assessing common cardiometabolic health biomarkers in blood samples [Time Frame: Baseline] Samples will be processed for the analysis of common biomarkers of cardiometabolic health: total cholesterol (mg/dL), HDL-cholesterol (mg/dL), triglycerides (mg/dL), glucose (mg/dL), insuline (uU/mL). Analyses will follow standardised routine procedures. • Assessing risk prediction scores [Time Frame: Baseline] Risk prediction scores (i.e., Framingham General Cardiovascular Risk Score, Framingham Heart Study Primary Risk Functions for heart disease, stroke, diabetes, fatty liver disease, and hypertension, Atherosclerotic Cardiovascular Disease (ASCVD) Risk, QRISK3®, QDScore®, Finnish Diabetes Risk Score (FINDRISC)) will be assessed to understand their relationship with the aggregate phenolic metabotypes observed. The higher the scores, the worse the risk of developing the disease. • Evaluating trimethylamine N-oxide (TMAO) in urine and plasma samples [Time Frame: Baseline] TMAO will be quantified in baseline urine and fasting plasma samples by UHPLC-MS/MS. • Evaluating eicosanoids in urine samples [Time Frame: Baseline] Eicosanoids, including prostaglandins, thromboxanes, leukotrienes, isoprostanes and neuroprostanes will be evaluated in baseline urine samples (0-h) by UHPLC- QqQ-MS/MS. • Assessing DNA oxidation catabolites and branched fatty acyl esters of hydroxyl fatty acids (FAHFAs) in plasma samples [Time Frame: Baseline] DNA oxidation catabolites and FAHFAs will be measured in fasting plasma samples by UHPLC-QqQ- MS/MS. • Determining genetic differences among subjects [Time Frame: Baseline] Genotyping will be conducted using genome wide, SNP array approach untargeted methodology, using commercially available SNP arrays and a tSNP approach. This approach will involve the genotyping of approximately 300 SNPs. Genomic DNA will be prepared from PBMCs by using	Actual Enrollment: Estimated Enrollment: 300 Original Estimated Enrollment: Same as current Age: 18 Years to 74 Years (Adult, Older Adult)	Study Sponsors: Same as current Collaborators: Azienda Ospedalier O- Universita ria di Parma University of Birmingha m Centro de Edafología y Biología Aplicada del Segura (CEBAS-	Study Start: May 31, 2022 Primary Completion: April 2023 (Final data collection date for primary outcome measure) Study Completion: April 2023 First Posted: June 10, 2022 Results First Posted: Last Update Posted: September 16,

	NCT Number	Title	Other Names	Status	Conditions	Interventions	Characteristics	Population	Sponsor/Collaborators	Dates
4	NCT0554 7360	Analysis of Blood	Title Acronym:	Recruiting	GastroIntestinal Bleeding	Not Provided	Study Type: Observational	Actual Enrollment:	Study Sponsors: Same as current	Study Start: July 19, 2022
		Metabolomics to Identify Potential	Other Ids: S18				Phase: Study Design: Observational Model: Cohort	Estimated Enrollment: 32	Collaborators: Not Provided	Primary Completion:
		Biomarkers of Gastrointestinal					Time Perspective: Cross-Sectional Primary Outcome Measures: Same as current	Original		December 31, 2022 (Final
		Bleeding Study					Secondary Outcome Measures: Not Provided	Estimated Enrollment: Same as current		data collection date for primary
		Documents:						Age: 21 Years		outcome measure)
								to 55 Years (Adult)		Study Completion:
								Sex: All		December 31, 2022
										First Posted: September 21, 2022
										Results First Posted:
										Last Update Posted: September 21, 2022
5	NCT0523	High-fat Meal	Title Acronym:	Active, not	Pediatric	Dietary Supplement: High-fat Challenge	Study Type: Interventional	Actual	Study Sponsors:	Study Start:
	0433	Challenge in Pediatrics	Other Ids: STUDY0200131	recruiting	Obesity • Insulin	The shake will be composed of a mixture of BOOST Glucose Control(R) (Nestlé Products) supplemented with	Phase: Not Applicable	Enrollment: Estimated	Same as current Collaborators:	May 1, 2022 Primary
		Study Documents:	6		Resistance	palm oil. Each participant will consume a volume of liquid equivalent to 25% of their estimated daily caloric needs, calculated by the USDA Dietary Reference Intakes using a moderate activity factor.	Study Design: Allocation: N/A Intervention Model: Single Group Assignment Masking: None (Open Label) Primary Purpose: Basic Science	Enrollment: 50 Original Estimated	Not Provided	Completion: September 19, 2022 (Final data collection
							Primary Outcome Measures: Same as current	Enrollment: Same as current		date for primary outcome
							Secondary Outcome Measures: Same as current	Age: 8 Years to 17 Years		measure)
								(Child)		Study Completion: December 31,
								Sex: All		2023
										First Posted: February 9, 2022
										Results First Posted:
										Last Update Posted: September 21, 2022

	NCT Number	Title	Other Names	Status	Conditions	Interventions	Characteristics	Population	Sponsor/Collaborat	Dates
6	NCT Number NCT0541 4409	Title The Gut Microbiome in Type 1 Diabetes and Mechanism of Metformin Action Study Documents:	Other Names Title Acronym: Other Ids: 15498	Not yet recruiting	Conditions • Type 1 Diabetes • Obesity	Interventions Drug: Metformin Metformin is an oral medication that improves insulin sensitivity.	Characteristics Study Type: Interventional Phase: Phase 2 Study Design: Allocation: Non-Randomized Intervention Model: Sequential Assignment Intervention Model Description: The investigators will examine the differences in the gut microbiome between lean and obese youth with type 1 diabetes and then enroll the obese youth with type 1 diabetes into the clinical trial to receive metformin as an intervention. Masking: None (Open Label) Primary Purpose: Treatment Primary Outcome Measures: • Differences in the gut microbiome in lean and obese youth with type 1 diabetes [Time Frame: 2 years] cross sectional comparison of stool microbiome using metagenomic sequencing data • Differences in the gut microbial metabolites in lean and obese youth with type 1 diabetes [Time Frame: 2 years] The investigators will measure and compare the stool and serum short chain fatty acids using mass spectrometry • Differences in the gut microbial metabolites in lean and obese youth with type 1 diabetes [Time Frame: 2 years] The investigators will measure and compare the stool and serum secondary bile acids using mass spectrometry • Changes in the gut microbiome in obese youth with type 1 diabetes in response to metformin [Time Frame: 3 years]	Actual Enrollment: Estimated Enrollment: 114 Original Estimated Enrollment: Same as current Age: 11 Years to 18 Years (Child, Adult) Sex: All	Sponsor/Collaborators Study Sponsors: Same as current Collaborators: Not Provided	Study Start: October 2022 Primary Completion: August 2025 (Final data collection date for primary outcome measure) Study Completion: August 2026 First Posted: June 10, 2022 Results First Posted: Last Update Posted: September 19, 2022
							obese youth with type 1 diabetes [Time Frame: 2 years] The investigators will measure and compare the stool and serum secondary bile acids using mass spectrometry • Changes in the gut microbiome in obese youth with type 1 diabetes in response to metformin [Time Frame: 3			

	NCT Number	Title	Other Names	Status	Conditions	Interventions	Characteristics	Population	Sponsor/Collaborat ors	Dates
7	NCT0554 3369	Study to Compare the Level of Elafibranor in Blood After Repeat Administration in Japanese and Non-Asian Healthy Participants Study Documents:	Title Acronym: Other Ids: CLIN-60190-450	Not yet recruiting	Healthy Volunteers	Drug: Elafibranor Oral Tablet Other Name: GFT505	Study Type: Interventional Phase: Phase 1 Study Design: Allocation: Non-Randomized Intervention Model: Parallel Assignment Masking: None (Open Label) Primary Purpose: Other Primary Outcome Measures: Same as current Secondary Outcome Measures: Same as current	Actual Enrollment: Estimated Enrollment: 48 Original Estimated Enrollment: Same as current Age: 18 Years to 55 Years (Adult) Sex: All	Study Sponsors: Same as current Collaborators: Not Provided	Study Start: September 15, 2022 Primary Completion: December 23, 2022 (Final data collection date for primary outcome measure) Study Completion: December 23, 2022 First Posted: September 16, 2022 Results First Posted: Last Update Posted: September 16, 2022
8	NCT0519 9610	An Open-Label, Single-Dose, Parallel-Group Study of the Pharmacokinetic s and Safety of EQ143 Study Documents:	Title Acronym: Other Ids: EQ143-102	Recruiting	Severe Hepatic Impairment	Drug: aumolertinib A single dose of 55-mg EQ143 tablet will be administered in the morning on Day 1, and participants will remain for 5 days (4 nights) in the study center for collection of blood samples and safety monitoring. Participants will attend outpatient follow-up visits on Days 5, 6, 8, and 9 for additional blood sampling and safety assessments. Other Name: HS-10296	Study Type: Interventional Phase: Phase 1 Study Design: Allocation: N/A Intervention Model: Single Group Assignment Masking: None (Open Label) Primary Purpose: Treatment Primary Outcome Measures: Same as current Secondary Outcome Measures: Same as current	Actual Enrollment: Estimated Enrollment: 12 Original Estimated Enrollment: Same as current Age: 18 Years to 75 Years (Adult, Older Adult) Sex: All	Study Sponsors: Same as current Collaborators: Not Provided	Study Start: March 22, 2022 Primary Completion: December 2022 (Final data collection date for primary outcome measure) Study Completion: December 2022 First Posted: January 20, 2022 Results First Posted: Last Update Posted: September 19, 2022

	NCT Number	Title	Other Names	Status	Conditions	Interventions	Characteristics	Population	Sponsor/Collaborators	Dates
9	NCT0504 2414	Acute Effects of Endurance Exercise on Breastmilk Composition Study Documents:	Title Acronym: Other Ids: 263493	Recruiting	Breastmilk	 Behavioral: High intensity interval training Four times four minutes treadmill interval training Behavioral: Moderate intensity training Moderate intensity treadmill training Behavioral: Resting No training 	Study Type: Interventional Phase: Not Applicable Study Design: Allocation: N/A Intervention Model: Sequential Assignment Masking: None (Open Label) Primary Purpose: Basic Science Primary Outcome Measures: Same as current Secondary Outcome Measures: Same as current	Actual Enrollment: Estimated Enrollment: 20 Original Estimated Enrollment: Same as current Age: 18 Years and older (Adult, Older Adult) Sex: Female	Study Sponsors: Same as current Collaborators: St. Olavs Hospital	Study Start: August 24, 2021 Primary Completion: June 1, 2025 (Final data collection date for primary outcome measure) Study Completion: June 1, 2025 First Posted: September 13, 2021 Results First Posted: Last Update Posted: September 19, 2022

	NCT Number	Title	Other Names	Status	Conditions	Interventions	Characteristics	Population	Sponsor/Collaborators	Dates
10	NCT0511 5513	Standardized Field Test for	Title Acronym:	Recruiting	Marijuana	Drug: Medium THC	Study Type: Interventional	Actual	Study Sponsors:	Study Start:
	3313	Marijuana	Other Ids: HHC- 2021-0333		Impairment	Marijuana flower with medium THC	Phase: Phase 1	Enrollment: Estimated	Same as current Collaborators:	August 25, 2022 Primary
		Impairment II Study Documents:	DTNH2216C000 22 (Other Identifier: Department of			Drug: Placebo THC Marijuana flower with no THC	Study Design: Allocation: Non-Randomized Intervention Model: Crossover Assignment Masking: Single (Participant) Primary Purpose: Other	Enrollment: 32 Original Estimated	 National Highway Traffic 	Completion: February 2024 (Final data collection date
			Transportation)				Primary Outcome Measures:	Enrollment: Same as current	Safety Administr ation	for primary outcome
							 Marijuana induced performance changes on Cogstate Groton Maze Learning task. [Time Frame: Post placebo 	Age: 18 Years	(NHTSA)	measure)
							at 30 min and post active dose administration at: 30, 60, 90, 120, 150, 180 min.]	to 55 Years (Adult)	Hartford Hospital	Study Completion:
							The Cogstate Groton Maze Learning task assesses spatial learning and memory, it will be administered prior to dosing and at various time points after dosing.	Sex: All	 National Institute on Drug 	First Posted: November 10,
							 Marijuana induced performance changes on Inquisit Maze Learning task. [Time Frame: Post placebo at 30 		Abuse (NIDA)	2021
							min and post active dose administration at: 30, 60, 90, 120, 150, 180 min.]			Results First Posted:
							The Inquisit Maze Learning task assesses spatial learning and memory, it will be administered prior to dosing and at various time points after dosing.			Last Update Posted: September 21,
							 Marijuana induced performance changes on the Time Reproduction Task. [Time Frame: Post placebo at 30 min and post active dose administration at: 30, 60, 90, 120, 150, 180 min.] 			2022
							The Time Reproduction Task assesses general motor coordination plus timing, it will be administered prior to dosing and at various time points after dosing.			
							Secondary Outcome Measures:			
							 Change in concentration of THC/metabolites in blood samples. [Time Frame: Post placebo at 5 min and post active dose administration at: 5, 30, 90, 150 min.] 			
							Blood samples with be collected at 5 times throughout each day to assess for changes of THC and its metabolite levels.			
							 Change in performance on simulated driving Road Tracking Task. [Time Frame: Post placebo at 10 min and post active dose administration at 210 min.] 			
							The Road Tracking Task measures operational control of the vehicle. Operational control is measured by standard deviation of lane position from the center point of the lane.			

	NCT Number	Title	Other Names	Status	Conditions	Interventions	Characteristics	Population	Sponsor/Collaborat	Dates
11	NCT0554 5501	Ketone Ester and Acute Salt (KEAS) in Young Adults Study Documents:	Title Acronym: Other Ids: AU IRB #22-025	Not yet recruiting	• Salt; Excess • Hypertensi on	 Dietary Supplement: No Salt, No -OHB Participants will consume the following for ten days. Enteric capsules will be filled with a dextrose placebo. The placebo supplement will be a -OHB- free, taste and viscosity-matched, beverage produced by KetoneAid. Dietary Supplement: High Salt, No -OHB Participants will consume the following for ten days. Enteric capsules will be filled with Morton's table salt. Sodium consumption will be normalized to caloric intake (2 mg Sodium/Calorie). The placebo supplement will be a -OHB-free, taste and viscosity- matched, beverage produced by KetoneAid. Dietary Supplement: High Salt, High -OHB Participants will consume the following for ten days. Enteric capsules will be filled with Morton's table salt. Sodium consumption will be normalized to caloric intake (2 mg Sodium/Calorie). Ketone beverage will be the -OHB supplement produced by KetoneAid. Participants will consume 24 mL (12 grams -OHB) of the ketone beverage three times a day (total 36 grams -OHB). 	Study Design: Allocation: Randomized Intervention Model: Crossover Assignment Intervention Model Description: Placebo-controlled, double-blinded, randomized Masking: Double (Participant, Investigator) Masking Description: Participants will be randomized to a condition order. A single lab member, not involved in data collection or analysis, will know condition order and contents and distribute them to participants. Primary Purpose: Basic Science Primary Outcome Measures: Same as current Secondary Outcome Measures: Same as current	Actual Enrollment: Estimated Enrollment: 35 Original Estimated Enrollment: Same as current Age: 18 Years to 39 Years (Adult) Sex: All	Study Sponsors: Same as current Collaborators: • University of Utah • University of Missouri-Columbia	Study Start: October 1, 2022 Primary Completion: September 30, 2025 (Final data collection date for primary outcome measure) Study Completion: September 30, 2026 First Posted: September 19, 2022 Results First Posted: Last Update Posted: September 19, 2022
12	NCT0551 5588	A Study in Healthy Men to Test How BI 690517 is Taken up and Handled by the Body Study Documents:	Title Acronym: Other Ids: 1378- 0013 2022-001818-18 (EudraCT Number)	Recruiting	Healthy	• Drug: BI 690517 (C-14) BI 690517 (C-14) • Drug: BI 690517 BI 690517	Study Type: Interventional Phase: Phase 1 Study Design: Allocation: Non-Randomized Intervention Model: Single Group Assignment Masking: None (Open Label) Primary Purpose: Treatment Primary Outcome Measures: Same as current Secondary Outcome Measures: Same as current	Actual Enrollment: Estimated Enrollment: 14 Original Estimated Enrollment: Same as current Age: 18 Years to 55 Years (Adult) Sex: Male	Study Sponsors: Same as current Collaborators: Not Provided	Study Start: September 19, 2022 Primary Completion: November 11, 2022 (Final data collection date for primary outcome measure) Study Completion: November 11, 2022 First Posted: August 25, 2022 Results First Posted: Last Update Posted: September 21, 2022

	NCT Number	Title	Other Names	Status	Conditions	Interventions	Characteristics	Population	Sponsor/Collaborat	Dates
13	NCT0554 7204	Impact of Coffeeberry Extract on Skill Performance During Simulated Match Play in Academy Football Players Study Documents:	Title Acronym: Other Ids: PEP- 2114	Recruiting	Skill Performan ce Soccer Performan ce Coffeeberr y Effects	Other: Beverage Free from added carbohydrates (contain 1g CHO only) and electrolytes are absent.	Study Type: Interventional Phase: Not Applicable Study Design: Allocation: Randomized Intervention Model: Crossover Assignment Masking: Quadruple (Participant, Care Provider, Investigator, Outcomes Assessor) Primary Purpose: Basic Science Primary Outcome Measures: Same as current Secondary Outcome Measures: Same as current	Actual Enrollment: Estimated Enrollment: 15 Original Estimated Enrollment: Same as current Age: 16 Years to 28 Years (Child, Adult) Sex: All	Study Sponsors: Same as current Collaborators: Not Provided	Study Start: June 30, 2022 Primary Completion: January 31, 2023 (Final data collection date for primary outcome measure) Study Completion: January 31, 2023 First Posted: September 21, 2022 Results First Posted: Last Update Posted: September 21, 2022
14	NCT0554 9622	Impact of Diet on the Gut-Muscle Axis in Older Adults Study Documents:	Title Acronym: Other Ids: GRANT1338686 5	Not yet recruiting	Sarcopenia Dietary Exposure Microbial Colonizati on	 Other: Low-soluble fiber diet 12 older adults (>65y) will be randomized to consume the USDA Guidelines for fiber intake (10g of total fiber/1000 calories), as the low-soluble fiber diet Other: High-soluble fiber diet 12 older adults (>65y) will be randomized to consume 34-35g of total fiber/1000 calories, as the high-soluble fiber diet 	Study Type: Interventional Phase: Not Applicable Study Design: Allocation: Randomized Intervention Model: Parallel Assignment Intervention Model Description: 24 older adults (> 65y) will be randomized to consume a high- or low-soluble fiber diet Masking: Quadruple (Participant, Care Provider, Investigator, Outcomes Assessor) Primary Purpose: Treatment Primary Outcome Measures: Same as current Secondary Outcome Measures: Same as current	Actual Enrollment: Estimated Enrollment: 24 Original Estimated Enrollment: Same as current Age: 65 Years and older (Older Adult) Sex: All	Study Sponsors: Same as current Collaborators: Not Provided	Study Start: September 15, 2022 Primary Completion: April 15, 2025 (Final data collection date for primary outcome measure) Study Completion: April 15, 2025 First Posted: September 22, 2022 Results First Posted: Last Update Posted: September 22, 2022

	NCT Number	Title	Other Names	Status	Conditions	Interventions	Characteristics	Population	Sponsor/Collaborators	Dates
15	NCT0440 7390	Effects of Nicotinamide Riboside on the Clinical Outcome of Covid-19 in the Elderly Study Documents:	Title Acronym: Other Ids: H- 20026601	Withdrawn	COVID	 Dietary Supplement: Nicotinamide riboside The patients will receive 1 g of nicotinamide riboside or placebo orally every morning for 14 days. Dietary Supplement: Placebo Placebo 	Study Type: Interventional Phase: Phase 2 Study Design: Allocation: Randomized Intervention Model: Parallel Assignment Intervention Model Description: Randomized double-blind case-control trial Masking: Quadruple (Participant, Care Provider, Investigator, Outcomes Assessor) Primary Purpose: Treatment Primary Outcome Measures: Same as current Secondary Outcome Measures: Same as current	Actual Enrollment: 0 Estimated Enrollment: Original Estimated Enrollment: 100 Age: 70 Years and older (Older Adult) Sex: All	Study Sponsors: Same as current Collaborators: Bispebjerg Hospital Elysium Health	Study Start: June 1, 2020 Primary Completion: May 1, 2021 (Final data collection date for primary outcome measure) Study Completion: May 1, 2022 First Posted: May 29, 2020 Results First Posted: Last Update Posted: September 22, 2022
16	NCT0507 0858	A Study to Examine the Efficacy and Safety of Pozelimab and Cemdisiran Combination Therapy in Adult Patients With Symptomatic Generalized Myasthenia Gravis Study Documents:	Title Acronym: Other Ids: R3918-MG-2018 2020-003272-41 (EudraCT Number)	Recruiting	Generalized Myasthenia Gravis	 Drug: Pozelimab + Cemdisiran Subcutaneous administration as described in the protocol Drug: Cemdisiran SC administration as described in the protocol Other Name: ALN-CC5 Other: Placebo SC administration as described in the protocol Drug: Pozelimab SC administration as described in the protocol Other Name: REGN3918 	Study Type: Interventional Phase: Phase 3 Study Design: Allocation: Randomized Intervention Model: Parallel Assignment Masking: Quadruple (Participant, Care Provider, Investigator, Outcomes Assessor) Primary Purpose: Treatment Primary Outcome Measures: Same as current Secondary Outcome Measures: Same as current	Actual Enrollment: Estimated Enrollment: 235 Original Estimated Enrollment: 210 Age: 18 Years and older (Adult, Older Adult) Sex: All	Study Sponsors: Same as current Collaborators: Not Provided	Study Start: December 14, 2021 Primary Completion: August 14, 2024 (Final data collection date for primary outcome measure) Study Completion: May 1, 2027 First Posted: October 7, 2021 Results First Posted: Last Update Posted: September 22, 2022

	NCT Number	Title	Other Names	Status	Conditions	Interventions	Characteristics	Population	Sponsor/Collabora	Dates
17	NCT0532 9142	ASSIST: A Surveillance Study of Illicit Substance Toxicity Study Documents:	Title Acronym: Other Ids: GN21AE239	Recruiting	Overdose, Drug Drug Use Drug Abuse Drug Toxicity Drug Effect Illicit Drug Use Illicit Drug Overdose Illicit Drug Intoxicatio n	Diagnostic Test: Surplus sample toxicology analysis Anonymised surplus blood sample will be analysed for drugs and their metabolites by way of Mass Spectrometry and LGC Group, Cambridge.	Study Type: Observational Phase: Study Design: Observational Model: Cohort Time Perspective: Prospective Primary Outcome Measures: Same as current Secondary Outcome Measures: Same as current	Actual Enrollment: Estimated Enrollment: 1000 Original Estimated Enrollment: Same as current Age: 16 Years and older (Child, Adult, Older Adult) Sex: All	Study Sponsors: Same as current Collaborators: Public Health Scotland	Study Start: August 19, 2022 Primary Completion: August 19, 2023 (Final data collection date for primary outcome measure) Study Completion: August 19, 2023 First Posted: April 14, 2022 Results First Posted: Last Update Posted: September 19, 2022
18	NCT0007 8078	Clinical and Laboratory Study of Methylmalonic Acidemia Study Documents:	Title Acronym: Other Ids: 040127 04-HG-0127	Recruiting	Organic Acidemia Methylmal onic Acidemia Inborn Errors of Metabolis m	Not Provided	Study Type: Observational Phase: Study Design: Observational Model: Cohort Time Perspective: Prospective Primary Outcome Measures: Not Provided Secondary Outcome Measures: Not Provided	Actual Enrollment: Estimated Enrollment: 2275 Original Estimated Enrollment: Age: 1 Month and older (Child, Adult, Older Adult) Sex: All	Study Sponsors: Same as current Collaborators: Not Provided	Study Start: June 7, 2004 Primary Completion: Not Provided Study Completion: Not Provided First Posted: February 19, 2004 Results First Posted: Last Update Posted: September 21, 2022

	NCT Number	Title	Other Names	Status	Conditions	Interventions	Characteristics	Population	Sponsor/Collaborators	Dates
19	NCT0538 6758	A Study to Evaluate Molnupiravir (MK-4482; MOV) in Participants With Severe Renal Impairment (MK-4482-003) Study Documents:	Title Acronym: Other Ids: 4482- 003 MK-4482-003 (Other Identifier: Merck)	Recruiting	Renal Impairment	Drug: Molnupiravir Four 200 mg capsules administered orally as a single dose Other Name: MK-4482; MOV; EIDD-2801	Study Type: Interventional Phase: Phase 1 Study Design: Allocation: Non-Randomized Intervention Model: Parallel Assignment Masking: None (Open Label) Primary Purpose: Treatment Primary Outcome Measures: Same as current Secondary Outcome Measures: Same as current	Actual Enrollment: Estimated Enrollment: 18 Original Estimated Enrollment: Same as current Age: 18 Years to 75 Years (Adult, Older Adult) Sex: All	Study Sponsors: Same as current Collaborators: Not Provided	Study Start: June 29, 2022 Primary Completion: November 7, 2022 (Final data collection date for primary outcome measure) Study Completion: November 18, 2022 First Posted: May 23, 2022 Results First Posted: Last Update Posted: September 16, 2022
20	NCT0324 4722	Maternal Metabolic and Molecular Changes Induced by Preconception Weight Loss and Their Effects on Birth Outcomes Study Documents:	Title Acronym: Other Ids: HUM00124673 1R01DK124862 (U.S. NIH Grant/Contract)	Recruiting	Obesity; Familial Pregnancy Related	 Dietary Supplement: Very-low energy Diet (VLED) Structured, intensive dietary intervention using liquid meal replacements aimed at providing 800 kcal/day with a weight loss goal of 15% from baseline Other: Standard of care (SOC) Standard consultation with registered dietitian to determine appropriate caloric deficit for a low calorie diet, education and advice to achieve weight loss in obese women. Standard of care for normal weight women 	Study Type: Interventional Phase: Not Applicable Study Design: Allocation: Randomized Intervention Model: Parallel Assignment Masking: None (Open Label) Primary Purpose: Basic Science Primary Outcome Measures: Rate of development of gestational diabetes [Time Frame: 40 weeks after conception or 24-28 weeks of gestation or 0 to 40 weeks after conception] The number of patients who develop gestational diabetes will be tracked throughout pregnancy. Diagnosis will be made based on one abnormal value per participant. Gestational diabetes will be measured using an oral glucose tolerance test. Secondary Outcome Measures: Birthweight [Time Frame: within 12 hours of birth] Newborn birthweight will be measured within 12 hours of birth.	Actual Enrollment: Estimated Enrollment: 352 Original Estimated Enrollment: 540 Age: 18 Years to 40 Years (Adult) Sex: Female	Study Sponsors: Same as current Collaborators: National Institute of Diabetes and Digestive and Kidney Diseases (NIDDK)	Study Start: April 10, 2018 Primary Completion: August 31, 2025 (Final data collection date for primary outcome measure) Study Completion: August 31, 2025 First Posted: August 9, 2017 Results First Posted: Last Update Posted: September 21, 2022

	NCT Number	Title	Other Names	Status	Conditions	Interventions	Characteristics	Population	Sponsor/Collaborat ors	Dates
21	NCT0555 1455	Endocrinological and Physiological Responses to Short-term Reduced Carbohydrate Availability in Males Study Documents:	Title Acronym: Other Ids: 22/SPS/026	Recruiting	 Energy Supply; Deficiency Carbohydr ate Availabilit y 	 Other: Nutritional/dietary intake manipulation ('Normal') Energy Intake provision (60 kcal/kg FFM/day) to elicit 'normal' energy availability (45 kcal/kg FFM/day), with 60% from carbohydrates. Other Name: Normal Carbohydrate Availability Other: Nutritional/dietary intake manipulation ('Low') Energy Intake provision (60 kcal/kg FFM/day), with 1.5 g/kg of carbohydrate and 70-80% fat intake, to elicit 'low' carbohydrate availability in energy balance (45 kcal/kg FFM/day). Other Name: Low Carbohydrate Availability 	Study Design: Allocation: Randomized Intervention Model: Crossover Assignment Intervention Model Description: Participants will complete two five-day trial-arms in randomised order, with an ~10-day washout period between interventions. Masking: None (Open Label) Masking Description: Masking not possible as food quantity/type will vary between interventions. Primary Purpose: Basic Science Primary Outcome Measures: Same as current Secondary Outcome Measures: Same as current	Actual Enrollment: Estimated Enrollment: 9 Original Estimated Enrollment: Same as current Age: 18 Years to 40 Years (Adult) Sex: Male	Study Sponsors: Same as current Collaborators: Not Provided	Study Start: May 4, 2022 Primary Completion: December 2022 (Final data collection date for primary outcome measure) Study Completion: December 2022 First Posted: September 22, 2022 Results First Posted: Last Update Posted: September 22, 2022
22	NCT0499 0869	Inflammation in COPD and the Effect of Nicotinamide Riboside Study Documents:	Title Acronym: Other Ids: NR-COPD	Completed	COPD	 Dietary Supplement: Nicotinamide Riboside The patients will receive 1 g of Nicotinamide Riboside or placebo orally every morning and evening for 6 weeks. Dietary Supplement: Placebo Placebo 	Study Type: Interventional Phase: Not Applicable Study Design: Allocation: Randomized Intervention Model: Parallel Assignment Intervention Model Description: Randomized, double-blind, placebo-controlled Masking: Quadruple (Participant, Care Provider, Investigator, Outcomes Assessor) Primary Purpose: Treatment Primary Outcome Measures: Same as current Secondary Outcome Measures: Same as current	Actual Enrollment: 60 Estimated Enrollment: Original Estimated Enrollment: Same as current Age: 60 Years and older (Adult, Older Adult) Sex: All	Study Sponsors: Same as current Collaborators: Bispebjerg Hospital Elysium Health	Study Start: July 1, 2021 Primary Completion: August 18, 2022 (Final data collection date for primary outcome measure) Study Completion: August 18, 2022 First Posted: August 5, 2021 Results First Posted: Last Update Posted: September 21, 2022

	NCT Number	Title	Other Names	Status	Conditions	Interventions	Characteristics	Population	Sponsor/Collaborat	Dates
23	NCT0279 5442	Daily Protein Intake Patterns on Energy Metabolism and the Motivation to Snack Study Documents:	Title Acronym: Other Ids: GFHNRC500	Recruiting	Obesity	 Other: Even protein 5 day intake of even protein 3 day rotating menu. Other: Skewed protein 5 day intake of skewed protein 3 day rotating menu. 	Phase: Not Applicable Study Design: Allocation: Randomized Intervention Model: Crossover Assignment Masking: None (Open Label) Primary Purpose: Other Primary Outcome Measures: Postprandial lipid oxidation rate [Time Frame: 4 hours after meal consumption] Secondary Outcome Measures: Relative reinforcing value (RRV) of energy-dense snack foods as assessed by indicator [Time Frame: Day 6] RRV of energy-dense snack foods will be assessed by evaluating the number of responses (mouse button presses) a participant is willing to complete to gain access to an energy- dense snack food or an alternative vegetable item.	Actual Enrollment: Estimated Enrollment: 40 Original Estimated Enrollment: Same as current Age: 20 Years to 60 Years (Adult) Sex: All	Study Sponsors: Same as current Collaborators: Not Provided	Study Start: June 2016 Primary Completion: September 2023 (Final data collection date for primary outcome measure) Study Completion: December 2023 First Posted: June 10, 2016 Results First Posted: Last Update Posted: September 19, 2022
24	NCT0554 5865	Chardonnay Marc and Vascular Response Study Documents:	Title Acronym: Other Ids: 1810396	Recruiting	Cardiovas cular Diseases Vascular Dilation Oxidative Stress	 Other: Low Flavanol Cocoa Powder Cocoa Powder providing 30 mg of cocoa flavanols Other: High Flavanol Cocoa Powder Cocoa Powder providing 435 mg of cocoa flavanols Other: Vine to Bar Chocolate - 2 servings Chocolate providing both cocoa flavanols and Chardonnay marc Other: Vine to Bar Chocolate - 1 serving Chocolate providing both cocoa flavanols and Chardonnay marc Other: Vine to Bar Chocolate covered almonds Almonds covered with Vine to Bar Chocolate that provides both cocoa flavanols and Chardonnay marc 	Study Type: Interventional Phase: Not Applicable Study Design: Allocation: Randomized Intervention Model: Crossover Assignment Intervention Model Description:	Actual Enrollment: Estimated Enrollment: 5 Original Estimated Enrollment: Same as current Age: 30 Years to 50 Years (Adult) Sex: Male	Study Sponsors: Same as current Collaborators: Not Provided	Study Start: September 1, 2022 Primary Completion: June 30, 2023 (Final data collection date for primary outcome measure) Study Completion: June 30, 2023 First Posted: September 19, 2022 Results First Posted: Last Update Posted: September 19, 2022

NCT Number Title	Other Names	Status	Conditions	Interventions	Characteristics	Population	Sponsor/Collaborators	t Dates
NCT0493 9883 Effects of Carvedilol on Cardiotoxicity in Cancer Patients Submitted to Anthracycline Therapy Study Documents:	Title Acronym: Other Ids: AVAP-NG 989	Recruiting	Cancer	 Drug: Carvedilol Carvedilol will be dispensed in a staggered and progressive manner, initially from 6.25 mg twice daily, then increased to 12.5 mg twice daily, until maximum dose of 25 mg twice daily or development of contraindications Drug: Placebo Patients will receive placebo in a presumed staggered and progressive manner similar to the intervention group. The placebo will ideally be maintained for up to 30 days after the end of chemotherapy. 	Phase: Phase 4 Study Design: Allocation: Randomized Intervention Model: Parallel Assignment Masking: Double (Participant, Investigator) Masking Description: Randomization will be in the proportion of 1: 1 (carvedilol x placebo). Both randomization and allocation of patients will be chosen in a veiled manner to patients and to assess. Data on randomization and allocation will be under custody of the Data analysis and safety committee. Primary Purpose: Prevention Primary Outcome Measures: Same as current Secondary Outcome Measures: Drop in ejection fraction within 24 months. [Time Frame: 24 months] Drop in ejection fraction greater than 10% and values less than 55% Reduction in myocardial strain in 24 months from the start of treatment. [Time Frame: 24 months] Relative reduction of more than 15% in myocardial strain Diastolic dysfunction within 24 months [Time Frame: 24 months] Development of diastolic dysfunction within 24 months Elevation of biomarkers during chemotherapy and up to 24 months of follow-up [Time Frame: 24 months] Elevation of biomarkers (NT-pro BNP and troponin) during chemotherapy and up to 24 months of follow-up [Time Frame: 24 months] Quality of life in up to 24 months. [Time Frame: 24 months] Quality of life measured by questionnaire in up to 24 months. Cardiovascular complications in 24 months. [Time Frame: 24 months] Cardiovascular complications (death, resuscitated cardiac arrest, myocardial infarction, heart failure and cardiac arrhythmias) in 24 months.	Actual Enrollment: Estimated Enrollment: 1018 Original Estimated Enrollment: Same as current Age: 18 Years and older (Adult, Older Adult) Sex: All	Study Sponsors: Same as current Collaborators: Ministério da Saúde	Study Start: August 1, 2021 Primary Completion: December 30, 2023 (Final data collection date for primary outcome measure) Study Completion: December 30, 2024 First Posted: June 25, 2021 Results First Posted: Last Update Posted: September 21, 2022

	NCT Number	Title	Other Names	Status	Conditions	Interventions	Characteristics	Population	Sponsor/Collaborators	t Dates
26	NCT0285 1511	Augmenting Cognitive Training In Older Adults Study Documents:	Title Acronym: Other Ids: IRB201600785- N-R R01AG054077 (U.S. NIH Grant/Contract) RF1AG071469 (U.S. NIH Grant/Contract) AWD00532 (Other Identifier: UFIRST)	Active, not recruiting	Aging	 Behavioral: Cognitive Training Cognitive Training employs an eight component, PositScience BrainHQ suite via its researcher portal. Device: tDCS (active stimulation) A Soterix Clinical Trials Direct Current Stimulator will apply 20 minutes of 2.0mA direct current through two biocarbon rubber electrodes encased in saline soaked 5cm x 7cm sponges (8cc of 0.9% saline solution per sponge) placed over the frontal cortices at F3 and F4 (10-20 system). Device: tDCS (sham stimulation) Sham stimulation is performed with the same device and all procedures will be identical except for the duration of stimulation. Participants will receive 30 seconds of 2 mA of direct current stimulation at the beginning of the session. Participants habituate to the sensation of tDCS within 30-60 seconds of stimulation. This procedure provides the same sensation of tDCS without the full duration of stimulation, making it a highly effective sham procedure. Behavioral: Educational Training Educational training involves watching educational videos produced by the National Geographic Channel, which cover a range of topics such as history, nature, and wildlife. Participants will be asked to complete questions on the content of the videos to ensure sustained attention. 	Study Type: Interventional Phase: Not Applicable Study Design: Allocation: Randomized Intervention Model: Parallel Assignment Masking: Double (Participant, Investigator) Primary Purpose: Treatment Primary Outcome Measures: • Improvements in CT-associated neurocognitive ability following CT and tDCS [Time Frame: Change from baseline to post assessment (3 months)] Composite measure of cognitive and functional abilities. • Improvements in neural functions following CT and tDCS [Time Frame: Change from baseline to post assessment (3 months)] activation in working memory, attentional brain systems, and other regions of interest (measure = beta values from SPM models) Secondary Outcome Measures: Not Provided	Actual Enrollment: 306 Estimated Enrollment: Original Estimated Enrollment: 360 Age: 65 Years to 89 Years (Older Adult) Sex: All	Study Sponsors: Same as current Collaborators: University of Arizona National Institute on Aging (NIA)	Study Start: August 8, 2017 Primary Completion: April 30, 2023 (Final data collection date for primary outcome measure) Study Completion: April 30, 2023 First Posted: August 1, 2016 Results First Posted: Last Update Posted: September 22, 2022