

Managed by **Duke** Clinical & Translational Science Institute

The MURDOCK Study Community Registry and Biorepository is a 12,526-participant community-based longitudinal cohort recruited from a 20-Zip Code region in the Southeastern United States (U.S.) that is centered in the city of Kannapolis, NC and encompasses Cabarrus County, NC.

Creation of the cohort was funded by a gift to Duke University from the David H. Murdock Institute for Business and Culture, with operational support from Duke's Clinical and Translational Science Award (CTSA) grant (UL1TR002553) and the Duke Clinical and Translational Science Institute (CTSI).

Consenting participants complete a baseline health questionnaire at enrollment, as well as a brief physical exam and collection of blood and urine. Consent includes permission to access to information from medical records, storage of collected samples in the biorepository, access to collected data and biospecimens for future approved research studies and contact regarding new research study opportunities.

Data have been organized into "storefronts" that summarize characteristics of a population of research interest as well as available data and samples for that population. The following sections summarize the sources of data in the MURDOCK Study database, as well as important descriptions and definitions to help understand the data presented in the "storefronts".

1 Participant self-reported data at baseline. The baseline questionnaire collects contact information, current residential street address, and primary physician; alternate contact information; date and place of birth; demographics; current or past diagnosis of 34 medical conditions; menopausal status in women; medications, vitamins and supplements; dietary and physical activity assessment; hours of sleep per night; tobacco and alcohol use; second-hand smoke exposure; and selected PROMIS® participant-reported outcomes domains. Socioeconomic data collected at baseline included marital status, highest level of education of participant and participant's mother and father, employment status, mother's and father's occupations, housing (type, how paid for, number of adults and children in the household) and total household income. In addition, a brief physical exam (vital signs, height, weight, and waist circumference) was conducted at enrollment.

Medical conditions: "Do you have, or have you ever had, any of the following [medical conditions]?" (yes, no, don't know). Counts are unique participants reporting yes to specific condition. Medications: "Please list any pharmaceutical and/or natural medications (including vitamins) that you are currently taking." Data are captured in free-text format as written by the participant and coded using RxNorm. Summary metrics are based on everything reported. Top 5 reported medications are limited to reported prescriptions.

2 Biorepository samples. Blood was collected at baseline and processed into the following specific samples: whole blood in EDTA for DNA extraction, whole blood in PAXgene for RNA extraction, plasma, serum and buffy coat in cryovials. Urine was collected and aliquoted in cryovials. Sample collection was not done systematically for MURDOCK enrollees; however, some nested sub cohorts and other studies enrolling MURDOCK registry participants include sample collection at follow up time points. All samples are stored at -80°C in a central biorepository current managed by Fisher BioServices, a division of Thermo Fisher Scientific, under a contractual agreement with Duke University.

Samples in inventory: Data are summarized by sample type as well as specific container and size. Participant counts are unique individuals with one ore more aliquots. Aliquot counts are all unique samples for a given type and container, size. Freezers is a calculation of approximate storage requirements based on sample type/size, box size, and number of boxes that can be stored per freezer.

3 Participant self-reported changes in health via annual follow up. Participants are asked to complete a follow-up form once a year around the time of their original enrollment date. Participants may update contact information, primary care physician/practice and alternate contact. PROMIS domains are repeated at each annual time point in order to capture changes in participant-reported outcomes over time. The form collects new incidence/diagnosis of the same 34 medical conditions surveyed at baseline. Hospitalizations during the past year are collected along with reason, as well as specific medical procedures. Participants may update their medication list to reflect current medications, vitamins and supplements being taken at the time of follow up form completion.

Vital status: Death reported by family member or alternate contact is confirmed by obituary as the primary source. Cause of death is not captured. Follow-up metrics: Follow-up is defined as complete if participant fills out the survey online or by mail or phone. Completeness is measured as surveys completed relative to years eligible to complete follow-up. Medical conditions: "Please indicate if you have received a new diagnosis of any of the following medical conditions in the past year (yes, no, don't know)". Counts and percentages are unique participants reporting yes to specific condition in follow-up for participants that did NOT report yes at baseline. Procedures: "Please indicate if you have any of the following medical procedures in the past year". Counts are unique participants reporting the specified procedure one or more times during follow up. Hospitalizations: Participants are asked to report if they have been hospitalized within the last year, for each hospitalization they are asked to list reason(s) for hospitalization, admission date and hospital name. Reasons for hospitalization are captured as free-text responses as written by participants. Responses are coded, when possible, in order to list the most frequently reported reasons for hospitalization. Medications: (see note above for medications reported at baseline). The denominator for data based on last follow-up are participants with at least one follow-up survey complete.

4 Electronic health record (EHR) data from regional healthcare providers. Duke has partnered with regional healthcare providers to integrate data from EHR systems for consented MURDOCK Study participants. Participants are identified in EHR systems with robust matching algorithms using common identifiers from the MURDOCK and EHR databases. Data are transferred under a data use agreement (DUA) with the specific provider organization which specifies the scope of data and frequency of transfers. Data availability vary by participant and depend on whether or not a participant has had one or more encounters with the healthcare provider system during the time period included in the dataset.

Available EHR datasets: Data are summarized by healthcare provider organizations. Counts are unique participants with one or more ICD codes in the EHR dataset. Available EHR domains: Data area summarized by domain in the EHR dataset. Counts are unique participants with one of more records (rows of data) for the specified domain. Insights from available EHR data: Specific EHR data related to the population of research interest is presented with granularity when possible.

**5** Additional data collection from studies with MURDOCK participants. MURDOCK Study participants may be recruited to enroll in additional research study opportunities by Duke researchers or other collaborators. Data sharing is a condition of collaboration with with the MURDOCK Study; therefore, data collected from MURDOCK Study participants and/or generated from biospecimens as part of additional research studies is returned for integration with all other MURDOCK registry data

"Storefronts" for nested sub-cohorts summarize surveys, assessments and/or other data collected specifically as part of enrollment and participation in the study. Samples in inventory: Samples are summarized if collected (see note above for samples collected at baseline). Participation in other studies: Counts are participants from the population of research interest enrolled in the specified study listed. Brief descriptions of relevant studies are listed along with a summary of study procedures and/or data collected.



Other autoimmune disease

Implantable cardiac defibrillator

Multiple sclerosis

Cervical cancer

Prostate cancer

MURDO	OCK Chronic Obst	ructive Pulmonary	Disease (COPE	) Observational Stu	udy, N=452					
Participant self-rep	orted characteristi	cs at MURDOCK S	tudy enrollmen	t (baseline, [Februa	ry 2009 - Ma	arch 2018	3])			
Demographics at baseline		Education at baseline								
Age		Baseline	Less than hig		48 (11%)					
Median (25th, 75th)	61 (53, 67)	High school g		141 (31%)						
Min, Max		37, 87	Some college		197 (44%)					
Sex			Bachelor's de		37 (8%)					
Female		235 (52%)	Master's or hi	gher professional de	29 (6%)					
Male		Income at baseline								
Race			Under \$10,00		66 (15%)					
American Indian & Alaska Native		2 (<1%)	\$10,000-29,9	123 (27%)						
Asian		1 (<1%) \$30,000-49,999				91 (20%				
Black or African American		100 (22%)	\$50,000-69,9	53 (12%)						
Native Hawaiian & Other Pacific	Islander	0	\$70,000-89,9	27 (6%)						
White/Caucasian		338 (75%)	\$90,000 or m	31 (7%)						
Other		3 (1%)	Don't know, n	61 (14%)						
Multiple		8 (2%)	Body mass i							
Don't know/Not sure/Not answere	ed	0	<18.5 (underv	10 (2%)						
Ethnicity		18.5 - 24.9 (n	105 (23%)							
Hispanic or Latino		9 (2%)	25 - 29.9 (overweight)				159 (35%)			
Non-Hispanic or Latino		436 (96%)	30+ (obese)	178 (39%)						
Don't know/Not sure/Not answere	Exercise at baseline									
Smoking history at baseline		Little to no ph	225 (50%)							
Smoked		443 (98%)	Weekend ligh	54 (12%)						
Never smoked		6 (1%)		Moderate activity 3x per week						
Don't know, no response	3 (<1%)	Heavy activity	•			129 (29%) 26 (6%)				
Current or prior medical condit	Heavy activity 5x per week 13 (3									
25 of 34 solicited medical condition	ons, listed by desce	ending frequency			onte at baco	lino	10 (070)			
High cholesterol		235 (52%)		, vitamins, supplem	ents at base	IIIIE	7 (2, 44)			
High blood pressure		234 (52%)	Median (25 <sup>th</sup> ,	7 (3, 11						
Depression		166 (37%)	10+ reported,		150 (33%)					
Emphysema or "COPD"	"COPD" 155 (34%)		Top 5 report	440 (040()						
Obesity	pesity		Albuterol				110 (24%)			
Osteoarthritis	steoarthritis		Lisinopril			98 (22%)				
Asthma		110 (24%)	Omeprazole				62 (14%)			
Diabetes		96 (21%)	Fluticasone		59 (13%)					
Thyroid disease		59 (13%)	Metformin 57 (13							
Rheumatoid arthritis		53 (12%)	Samples currently in inventory (collected at baseline time point)							
Osteoporosis/Osteopenia		52 (12%)	Sample	Container, Size	Participants	Aliquots	Freezers			
Other mental illness		49 (11%)	Plasma	Cryovial, 0.5 mL	261	2,356	0.042			
Skin cancer, not melanoma		45 (10%)	Serum	Cryovial, 0.5 mL	260	1,535	0.027			
Coronary artery disease		41 (9%)		Cryovial, 5.0 mL	236	236	0.008			
Heart attack or angina		37 (8%)	Whole blood	PAXgene RNA	194	295	0.017			
Gout		36 (8%)		Vacutainer, 2.0 mL		191	0.006			
Atrial fibrillation		33 (7%)	Buffy coat	Cryovial, 2.0 mL	0	0	0.000			
Stroke		28 (6%)	Urine	Cryovial, 10.0 mL	243	243	0.019			
Other type of cancer		26 (6%)	Total			4,856	0.119			
Congestive heart failure		25 (6%)								
		ì								

24 (5%)

15 (3%)

14 (3%) 12 (3%)

11 (2%)

72 (16%)



Lipid panel

## MURDOCK Chronic Obstructive Pulmonary Disease (COPD) Observational Study, N=452

	Participant state	is and da	ta from MU	IBDOCK 6	udy follow-up surveys and electronic health reco	rds				
Dorticin -	•	is and ud	ta iroiii iviO	MDOOK 3	New medical condition diagnoses reported i		n			
Participant vital status				(	16 of 34 solicited medical conditions, listed by a					
	Alive			381 (84%	Emphysema or "COPD"	95 / 297 (32%)				
Deceased				71 (16%	Osteoarthritis	81 / 335 (24%				
Current Age				Currer	High blood pressure	69 / 218 (32%)				
Median (25 <sup>th</sup> , 75 <sup>th</sup> )				70 (61, 77	High cholesterol	67 / 217 (31%)				
Min, Max				46, 90	Rheumatoid arthritis	60 / 399 (15%)				
	p metrics, study participa				Skin cancer, not melanoma	56 / 407 (14%)				
Median (25th, 75th) months since enrollment			1	17 (77, 159	Thyroid disease		49 / 393 (12%)			
`	25th, 75th) years since enrolli			10 (7, 14	Obesity		/ 333 (14%)			
Median (2	25 <sup>th</sup> , 75 <sup>th</sup> ) yearly follow-ups	complete		6 (3, 10	Osteoporosis/Osteopenia	45 / 400 (11%)				
Overall co	ompleteness of follow-up, n	/N (%)	2,907/	3,652 (80%	) Depression	43 / 286 (15%)				
At least o	ne (1) follow-up survey com	plete, n (%	6)	426 (94%	Asthma	41 / 342 (12%)				
100% cor	npletion (n, %)			190 (42%	) Diabetes	39 / 356 (11%)				
Last comp	oleted follow-up ≤ 18 month	s		200 (44%	Coronary artery disease	35 / 411 (9%)				
Enrolled i	n one or more other studies			452 (100%	Congestive heart failure	34 / 427 (8%)				
Available	EHR datasets by source	(any ICD	code)		Atrial fibrillation	34 / 419 (8%)				
Any source	e			137 (30%	Other mental illness	32 / 403 (8%)				
Novant H	ealth			80 (18%			27 100 (070)			
Cabarrus Health Alliance				57 (13%			054 (700()			
Cabarrus Rowan Community Health Centers			24 (5%)			351 (78%)				
Bethesda Health Center					Chest x-ray	332 (73%)				
Communit	ty Free Clinic		5 (1%)			253 (56%)				
Atrium (Carolinas Healthcare)					Heart/cardiac stress test	184 (41%)				
Available EHR data domains					Joint replacement	62 (14%)				
Diagnoses				137 (30%	Heart/cardiac catheterization	59 (13%) 57 (13%)				
Labs				94 (21%	Heart/cardiac angioplasty or stent		` ′			
Vitals				76 (17%			19 (4%)			
Medications				101 (22%			220 (520()			
Allergies				42 (9%	Participants reporting 1 or more hospitalizations		239 (53%)			
Immunizations				32 (7%	Unique hospitalizations reported	398				
Problems				64 (14%	Median (25th, 75th) hospitalizations reported		2 (1, 3)			
Procedures				52 (12%	Coded reasons for self-reported hospitalization listed in descending frequency	Events	Participants			
Hospitalizations				40 (9%	Uncoded	305	163			
Insights from available EHR data					Pneumonia	47	34			
Date range: Sep. 1993 (first encounter), Aug. 2022 (last encounter)					Surgery	35	33			
Median (25 <sup>th</sup> , 75 <sup>th</sup> )				3 (104, 327		26	22			
,	necodes, mapped from dia	anosis c		. , . ,	Chest Pain	22				
Phecode	Description	Group		n, ppts			17			
272.1	Hyperlipidemia		e/metabolic		Body mass index (BMI) at most recent comp	leted folio				
401.1	Essential hypertension		ry system	31	<18.5 (underweight)	15 (4%)				
250.2	Type 2 diabetes		e/metabolic		18.5 - 24.9 (normal weight)	106 (25%)				
530.1	Esophagitis, GERD	Digestive			25 - 29.9 (overweight)	137 (32%)				
296.2	Depression		disorders 12		30+	168 (39%)				
512.8	Cough	Respirat			Medications, vitamins, supplements at most	recent follow up				
Select laboratory tests					Median (25th, 75th) reported					
Test			Labs Participants		10+ reported, n (%)		156 (35%)			
Comprehensive metabolic panel			357	4	Ton E reported medications					
CBC and differential			269	4	Atomicototic	113 (25%)				
Basic metabolic panel			255	4		94 (21%)				
TSH			196	3			81 (18%)			
Hemoglobin A1c			205	3		77 (17%)				
			200	-	Louisopili		(,0)			

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Metoprolol



## MURDOCK COPD Observational Study, study design and assessments

Full protocol title: MURDOCK COPD Observational Study, the relationship between GOLD risk group and clinical outcomes in a community-based COPD cohort

Study investigators

Principal investigator: Scott Palmer, MD, MHS Co-principal investigator: Jamie Todd, MD

Study phenotypes

Met COPD criteria: 254

Met SRS or PRISM criteria: 198

Met SRS criteria only: 113 Met PRISM criteria only: 76

Met both SRS and PRISM criteria: 9

Study definitions

COPD: FEV1/FVC ratio, measured by spirometry, < 0.70

FEV1: Forced expiratory volume in one second

FVC: Forced vital capacity, total amount of air exhaled during an

FEV test

SRS: Symptomatic smoker with respiratory symptoms, FEV1/FVC >=

0.70 AND FVC >= 80% of predicted AND CAT score of >= 10

CAT: COPD assessment test

Preserved ratio impaired spirometry (PRISm), FEV1/FVC >= 0.70

AND FEV1 < 80% of predicted

GOLD: Global initiative for Chronic Obstructive Lung Disease

The study schedule of assessments is included below. The study was discontinued by the Sponsor during study month 12 assessments. A critical variables report of data from baseline and available follow-up time points was generated. The study investigators should be contacted regarding these data.

Visit Number <sup>1</sup>	Pre / Screening Visit <sup>2</sup>	Enrollment Visit	Follow-Up Visits								Early Term		
	Visit 0	Visit 1	Visit2	Visit 3	Visit 4	Visit 5	Visit 6	Visit7	Visit 8	Visit 9	Visit 10	Visit 11	
Study Month	0	0	6	12	18	24	30	36	42	48	54	60	
Informed Consent	X												
Medical record release and HIPAA form		Х		х		Х		Х		х		Х	x
Demographic data	X	Х											
Medical history <sup>3</sup>	X	X		Χ		Χ		Χ		Х		Х	
Exacerbation history <sup>3</sup>		X	Χ	Χ	Χ	Χ	Χ	Χ	Χ	Χ	Χ	Χ	
Concomitant medication <sup>4</sup>		Х	Χ	Χ	Χ	Χ	Х	Х	Χ	Х	Х	Х	
Self-reported hospitalizations		Х	Х	Х	Х	Х	Х	Х	Х	Х	Х	Х	
Hospital bill/medical record collection <sup>5</sup>		х	Х	х	Х	Х	х	х	Х	х	Х	Х	
Safety event collection and reporting <sup>6</sup>		х	Х	Х	X	X	Х	Х	Х	Х	Х	Х	
Confirm vital status			Χ	Χ	Χ	Χ	Χ	Χ	Χ	Х	Χ	Х	X
COPD assessment test	X			Χ		Χ		Χ		Χ		Х	
IPAQ-Short Form		X	Χ	Χ	Χ	Χ	Χ	Х	Χ	Χ	Χ	Χ	
Six-minute walk test		X											
Height and weight	X			Х		X		X		Х		Χ	
Spirometry	X			Х		Х		Х		Χ		Х	
Assign GOLD risk group <sup>7</sup>		X		Х		Χ		Х		Χ		Х	
Document termination reason <sup>8</sup>													Х

1 Visits 2, 4, 6, 8 and 10 will be conducted via standardized telephone interview at the six-month interval between annual in-person study visits. A standardized interview script will be used to elicit patient-reported information.

2 At prescreening, verbal consent will be obtained. Contact information will be collected or confirmed (for subjects already in the MURDOCK Registry), a subject number will be assigned, and age, smoking history, and previous/current lung transplant listing status will be collected.

3 A brief medical history review will be completed at prescreening to determine the subject's smoking history. If the subject is deemed to be eligible after all screening procedures are completed, then a detailed medical history and exacerbation history review include determination of the subject's burden of respiratory exacerbations within the past one year, and common COPD comorbidities will be completed. A brief medical history review including interval exacerbations will be updated at each annual assessment to capture interval changes in self-reported health status.

4 Concomitant medications recorded should include all prescription medications (including short-acting medications/inhalers, maintenance medications/inhalers, rescue medications/inhalers, antibiotics, oxygen, and any other medications taken for COPD or COPD comorbidities). Routine over-the-counter medication use (ex. Advil, Tylenol) does not need to be collected.

5 The hospital bill and discharge summary will be collected for self-reported hospitalizations; confirmation of the hospitalization, date of admission, date of discharge, discharge medications (if available), and ICD-9 or 10 codes for primary and secondary diagnoses will be entered into the database. The hospital bill will be the primary source of information for hospitalization confirmation, date of admission, date of discharge, and ICD-9 or 10 codes. The discharge summary will be the primary data source for the discharge medications.

6 See Section 5: Safety Event Reporting and Follow-Up for more detail on event collection and reporting to BI.

7 GOLD risk group will be assigned (if applicable) using a computer-based algorithm following the study visits.

8 If a subject terminates early from the study, indicate the date and reason for withdrawal in the database