DEPARTMENT OF HEALTH AND HUMAN SERVICES CENTERS FOR MEDICARE & MEDICAID SERVICES

Form Approved OMB No. 0938-0581

CLINICAL LABORATORY IMPROVEMENT AMENDMENTS (CLIA) APPLICATION FOR CERTIFICATION

I. GENERAL INFORMATION						
☐ Initial Application ☐ Survey			CLIA IDENTIFICATION NUMBER			
Change in Certificate Type			51	587		
X Other Changes (Specify) Director						
F//			(If an initial application leave blan	k, a number will b	e assigned)	
Effective Date						
FACILITY NAME			FEDERAL TAX IDENTIFICATION NUMBER			
One Really Great Lab, LLC			550123456			
EMAIL ADDRESS OneReallyGreatLab@dmail.com			TELEPHONE NO. (Include area code) 304-555-1234	FAX NO. (Include area code) 304-555-2232		
FACILITY ADDRESS — Physical Location of Laboratory (Building, Floor, Suite if applicable.) Fee Coupon/Certificate will be mailed to this Address unless mailing or corporate address is specified			MAILING/BILLING ADDRESS (If different from facility address) send Fee Coupon or certificate			
NUMBER, STREET (No P.O. Boxes)			NUMBER, STREET			
2037 Success Street						
CITY Best City	STATE	ZIP CODE 25163	CITY	STATE	ZIP CODE	
	SEND CERTIFICATE	A CONTROL OF	CORPORATE ADDRESS (If different f	rom facility) send Fee	Coupon or certificate	
X Physical	× Physical		NUMBER, STREET			
Mailing	Mailing					
Corporate	Corporate					
NAME OF DIRECTOR (Last, First, Midd			CITY	STATE	ZIP CODE	
Stokes, Justin L.						
CREDENTIALS			FOR OFFICE USE ONLY			
M.D.			Date Received			
II. TYPE OF CERTIFICATE REC certificate testing requirements		k only one) Plea	se refer to the accompanying i	nstructions for in	nspection and	
☐ Certificate of Waiver (Complete Sections I – VI and IX – X)						
Certificate for Provider Performed Microscopy Procedures (PPM) ((Complete Sections I-VII and IX-X)						
▼ Certificate of Compliance (Complete Sections I – X)						
☐ Certificate of Accreditation (Complete Sections I – X) and indicate which of the following organization(s) your laboratory is accredited by for CLIA purposes, or for which you have applied for accreditation for CLIA purposes.						
☐ The Joint Commission ☐ AOA ☐ AABB ☐ A2LA						
☐ CAP	co	DLA 🗌	ASHI			
If you are applying for a Certificate of Accreditation, you must provide evidence of accreditation for your laboratory by an approved accreditation organization as listed above for CLIA purposes or evidence of application for such accreditation within 11 months after receipt of your Certificate of Registration.						

NOTE: Laboratory directors performing non-waived testing (including PPM) must meet specific education, training and experience under subpart M of the CLIA regulations. Proof of these qualifications for the laboratory director must be submitted with this application.

PRA Disclosure Statement

According to the Paperwork Reduction Act of 1995, no persons are required to respond to a collection of information unless it displays a valid OMB control number. The valid OMB control number for this information collection is 0938-0581. Expiration Date: 3/31/2021. The time required to complete this information collection is estimated to average one hour per response, including the time to review instructions, search existing data resources, gather the data needed, and complete and review the information collection. If you have comments concerning the accuracy of the time estimate(s) or suggestions for improving this form, please write to: CMS, 7500 Security Boulevard, Attn: PRA Reports Clearance Officer, Mail Stop C4-26-05, Baltimore, Maryland 21244-1850. *****CMS Disclaimer*****Please do not send applications, claims, payments, medical records or any documents containing sensitive information to the PRA Reports Clearance Office. Please note that any correspondence not pertaining to the information collection burden approved under the associated OMB control number listed on this form will not be reviewed, forwarded, or retained. If you have questions or concerns regarding where to submit your documents, please contact LabExcellence@cms.hhs.gov.

III.	TYPE OF L	ABORATORY (Check the one mo	st descriptive of fa	cility type)			
□ 01 Ambulance □ 11 Health Main. Organizati □ 02 Ambulatory Surgery Center □ 12 Home Health Agency □ 03 Ancillary Testing Site in □ Health Care Facility □ 13 Hospice □ 04 Assisted Living Facility □ 15 Independent □ 05 Blood Bank □ 16 Industrial □ 06 Community Clinic □ 17 Insurance □ 07 Comp. Outpatient Rehab Facility □ 18 Intermediate Care Facility □ 08 End Stage Renal Disease □ Dialysis Facility □ 19 Mobile Laboratory □ 10 Health Fair □ 20 Pharmacy □ 10 Health Fair □ 21 Physician Office			Agency Care Facilities for h Intellectual tory	22 Practitioner Other (Specify) 23 Prison 24 Public Health Laboratories 25 Rural Health Clinic 26 School/Student Health Service 27 Skilled Nursing Facility/				
IV.	HOURS OF	LABORATORY	TESTING (List til	mes during which lab	ooratory testing is pe	erformed in HH:MM	format) If testing 2	24/7 Check Here 🗵
		SUNDAY	MONDAY	TUESDAY	WEDNESDAY	THURSDAY	FRIDAY	SATURDAY
	FROM:							
	TO:							
(For	multiple sites,	attach the additi	onal information	using the same for	mat.)			
٧.	MULTIPLE S	ITES (must meet	one of the regula	tory exceptions to	apply for this pro	vision in 1-3 belov	v)	
Ar	you applyir	ng for a single si	ite CLIA certifica	te to cover mult	tiple testing loca	tions?		1
X	No. If no, go	to section VI.	☐ Yes. If yes	, complete rema	inder of this sec	tion.		
1.	 Indicate which of the following regulatory exceptions applies to your facility's operation. Is this a laboratory that is not at a fixed location, that is, a laboratory that moves from testing site to testing site, such as mobile unit providing laboratory testing, health screening fairs, or other temporary testing locations, and may be covered under the certificate of the designated primary site or home base, using its address? Yes No If yes and a mobile unit is providing the laboratory testing, record the vehicle identification number(s) (VINs) and attach to the application. 					ay be covered) and attach to the		
2.	Is this a not-for-profit or Federal, State or local government laboratory engaged in limited (not more than a combination of 1 moderate complexity or waived tests per certificate) public health testing and filing for a single certificate for multiple sites? Yes \sum No					or		
	If yes, provide the number of sites under the certificate and list name, address and test performed for each							
3.	site below. Is this a hospital with several laboratories located at contiguous buildings on the same campus within the same physical location or street address and under common direction that is filing for a single certificate for these locations? Yes No							
If yes, provide the number of sites under this certificate and list name or department, location within hospital and specialty/subspecialty areas performed at each site below.								
	If additional	space is neede	d, check here 🗌	and attach the	additional infor	mation using the	e same format.	
NAME AND ADDRESS/LOCATION NAME OF LABORATORY OR HOSPITAL DEPARTMENT			1	ESTS PERFORMI	D/SPECIALTY/S	UBSPECIALTY		
NA	ME OF LABORAT	ORY OR HOSPITAL L	DEPARTMENT					
AD	DRESS/LOCATION	N (Number, Street, Lo	ocation if applicable)					
CIT	Y, STATE, ZIP CO	DE	TELEPHON	E NO. (Include area c	ode)			
NA	ME OF LABORAT	ORY OR HOSPITAL D	DEPARTMENT					
ADDRESS/LOCATION (Number, Street, Location if applicable)								
CITY, STATE, ZIP CODE TELEPHONE NO. (Include area code)			ode)					

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In the next three sections, indicate testing performed and annual test volume.
VI. WAIVED TESTING If only applying for a Certificate of Waiver, complete this section and skip sections VII (PPM Testing) and VIII (Non-Waived Testing).
dentify the waived testing (to be) performed. Be as specific as possible. This includes each analyte test system or device used the laboratory. e.g. (Rapid Strep, Acme Home Glucose Meter)
cme Rapid Strep
cme Urine Dipsticks
Indicate the ESTIMATED TOTAL ANNUAL TEST volume for all waived tests performed 400
☐ Check if no waived tests are performed
f additional space is needed, check here \square and attach additional information using the same format.
VII. PPM TESTING If only applying for a Certificate for PPM, complete this section and skip section VIII (Non-Waived Testing).
Identify the PPM testing (to be) performed. Be as specific as possible. e.g. (Potassium Hydroxide (KOH) Preps, Urine Sediment Examinations)
Indicate the ESTIMATED TOTAL ANNUAL TEST volume for all PPM tests performed
If also performing waived complexity tests, complete Section VI. For laboratories applying for certificate of compliance or certificate of accreditation, also include PPM test volume in the specialty/subspecialty category and the "total estimated an test volume" in section VIII.
Check if no PPM tests are performed
If additional space is needed, check here \square and attach additional information using the same format.

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VIII. NON-WAIVED TESTING (Including PPM testing if applying for a Certificate of Compliance or Accreditation) Complete this section only if you are applying for a Certificate of Compliance or a Certificate of Accreditation.

Identify the non-waived testing (to be) performed. Be as specific as possible. This includes each analyte test system or device used in the laboratory e.g. (Potassium, Acme Chemistry Analyzer).

Acme Blood Chemistry Analyzer Acme CBC

If additional space is needed, check here \square and attach additional information using the same format.	

If you perform testing other than or in addition to waived tests, complete the information below. If applying for one certificate for multiple sites, the total volume should include testing for ALL sites.

Place a check (/) in the box preceding each specialty/subspecialty in which the laboratory performs testing. Enter the estimated annual test volume for each specialty. Do not include testing not subject to CLIA, waived tests, or tests run for quality control, calculations, quality assurance or proficiency testing when calculating test volume. (For additional guidance on counting test volume, see the instructions included with the application package.)

If applying for a Certificate of Accreditation, indicate the name of the Accreditation Organization beside the applicable specialty/ subspecialty for which you are accredited for CLIA compliance. (The Joint Commission, AOA, AABB, A2LA, CAP, COLA or ASHI)

SPECIALTY / SUBSPECIALTY	ACCREDITING ORGANIZATION	ANNUAL TEST VOLUME	SPECIALTY / SUBSPECIALTY	ACCREDITING ORGANIZATION	ANNUAL TEST VOLUME
HISTOCOMPATIBILITY 010			HEMATOLOGY 400		4000
Transplant			★ Hematology		
Nontransplant			IMMUNOHEMATOLOGY		
MICROBIOLOGY			ABO Group & Rh Group 510		
Bacteriology 110			Antibody Detection (transfusion) 520		
☐ Mycobacteriology 115			Antibody Detection (nontransfusion) 530		
☐ Mycology 120			Antibody Identification 540		
Parasitology 130			Compatibility Testing 550		
☐ Virology 140			PATHOLOGY		
DIAGNOSTIC IMMUNOLOGY			Histopathology 610		
Syphilis Serology 210			Oral Pathology 620		
General Immunology 220			Cytology 630		
CHEMISTRY 3000		3000	RADIOBIOASSAY 800		
■ Routine 310			Radiobioassay		
Urinalysis 320			CLINICAL CYTOGENETICS 900		
Endocrinology 330			Clinical Cytogenetics		
☐ Toxicology 340			TOTAL ESTIMATED ANNUA	7000	

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IX. TYPE OF CONTROL (check the one most descriptive of ownership type)					
VOLUNTARY NONPROFIT	FOR PROFIT	GOVERNMENT			
□ 01 Religious Affiliation	□ 04 Proprietary □	□ 05 City			
□ 02 Private Nonprofit		☐ 06 County			
\square 03 Other Nonprofit		□ 07 State			
		□ 08 Federal			
(Specify)		\square 09 Other Government			
		(Specify)			
V DIDECTOR ASSULATION WITH OTHER		(specify)			
X. DIRECTOR AFFILIATION WITH OTHI	ER LABORATORIES				
If the director of this laboratory serves as director for additional laboratories that are separately certified, please complete the following:					
CLIA NUMBER	NAME OF LABORATORY				
51D1234566	Another Gr	reat Lab, Inc.			
ATTENTION: READ THE FOLLOWING CAREFULLY BEFORE SIGNING APPLICATION					
Any person who intentionally violates or any regulation promulgated thereu 18, United States Code or both, except requirement such person shall be imprunited States Code or both.	nder shall be imprisoned for not more that if the conviction is for a second c	than 1 year or fined under title or subsequent violation of such a			
Consent: The applicant hereby agrees to applicable standards found necessary by section 353 of the Public Health Service any Federal officer or employee duly of its pertinent records at any reasonable determine the laboratory's eligibility of requirements.	by the Secretary of Health and Human e Act as amended. The applicant further lesignated by the Secretary, to inspect time and to furnish any requested information or treatment of the continued eligibility for its certificate	Services to carry out the purposes of er agrees to permit the Secretary, or the laboratory and its operations and ormation or materials necessary to			
PRINT NAME OF OWNER/DIRECTOR OF LABORATORY					
Justin L. Stokes SIGNATURE OF OWNER/DIRECTOR OF LABORAT	TORY (Sign in ink)	DATE 07-19-2018			
NOTE: Completed 116 applications must be sent to your local State Agency. Do not send any payment with your completed 116 application.					

STATE AGENCY CONTACT INFORMATION CAN BE FOUND AT:

http://www.cms.gov/Regulations- and - Guidance/Legislation/CLIA/Downloads/CLIASA.pdf