

Fermentative Production of L-2-Hydroxyglutarate by Engineered *Corynebacterium glutamicum* via Pathway Extension of L-Lysine Biosynthesis

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Prell C, Burgardt A, Meyer F and Wendisch VF (2021) Fermentative Production of L-2-Hydroxyglutarate by Engineered Corynebacterium glutamicum via Pathway Extension of L-Lysine Biosynthesis. Front. Bioeng. Biotechnol. 8:630476. doi: 10.3389/fbioe.2020.630476 L-2-hydroxyglutarate (L-2HG) is a trifunctional building block and highly attractive for the chemical and pharmaceutical industries. The natural L-lysine biosynthesis pathway of the amino acid producer Corynebacterium glutamicum was extended for the fermentative production of L-2HG. Since L-2HG is not native to the metabolism of C. glutamicum metabolic engineering of a genome-streamlined L-lysine overproducing strain was required to enable the conversion of L-lysine to L-2HG in a six-step synthetic pathway. To this end, L-lysine decarboxylase was cascaded with two transamination reactions, two NAD(P)-dependent oxidation reactions and the terminal 2-oxoglutarate-dependent glutarate hydroxylase. Of three sources for glutarate hydroxylase the metalloenzyme CsiD from Pseudomonas putida supported L-2HG production to the highest titers. Genetic experiments suggested a role of succinate exporter SucE for export of L-2HG and improving expression of its gene by chromosomal exchange of its native promoter improved L-2HG production. The availability of Fe²⁺ as cofactor of CsiD was identified as a major bottleneck in the conversion of glutarate to L-2HG. As consequence of strain engineering and media adaptation product titers of $34 \pm 0 \,\text{mM}$ were obtained in a microcultivation system. The glucose-based process was stable in 2L bioreactor cultivations and a L-2HG titer of 3.5 g L^{-1} was obtained at the higher of two tested aeration levels. Production of L-2HG from a sidestream of the starch industry as renewable substrate was demonstrated. To the best of our knowledge, this study is the first description of fermentative production of L-2HG, a monomeric precursor used in electrochromic polyamides, to cross-link polyamides or to increase their biodegradability.

Keywords: C. glutamicum, L-2-hydroxyglutarate, metabolic engineering, glutarate hydroxylase, wheat sidestream concentrate, bioreactor

INTRODUCTION

L-2-hydroxyglutarate (L-2HG) can be obtained by hydroxylation of the C₅-dicarboxylic acid glutarate at the α -carbon position. Glutarate is known to be a demanded building block for the production of biopolymers, like aliphatic polyamide 6,5 which is mainly used in the construction industry (Navarro et al., 1997). Carbon chain length and functional groups of the monomers

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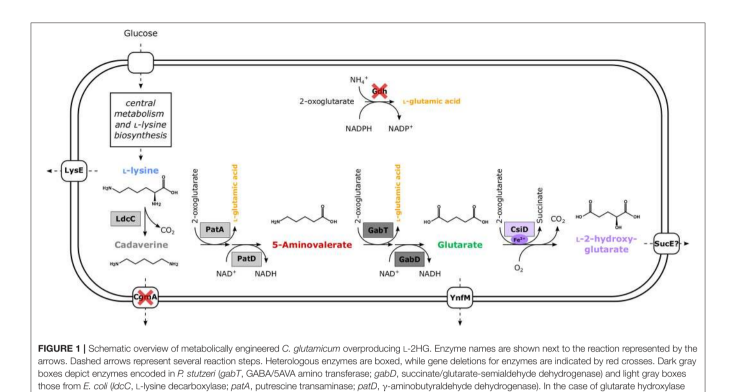
are important for the performance of the biopolymer with respect to physical and chemical properties. In the case of L-2HG, the additional hydroxyl group provides greater versatility for further chemical modification with the potential to alter the properties of the resulting polymer. Furthermore, incorporation of hydroxylated monomers enhances the biodegradability of the polyamide (Varela and Orgueira, 2000). Production of L-2HG by microbial fermentation would provide the chemical industry with an environmentally friendly building block for polyamides and polyesters. The addition of functionalities on polyester backbones (hydroxyl, carboxyl, allyl, azido, and acetylene groups) facilitates covalent post-polymerization modification. In 2017, it was shown that the enantiomer D-2-hydroxyglutarate (D-2HG) can be used as a functionalized building block for a polyester after cyclization to an allyl containing lactide (Nan and Feher, 2017). Also, it was demonstrated that electrochromic polyamides with functional hydroxyl groups for homogeneous hybrid films can be produced. The repeating units of hydroxysuccinate in the polymer backbone provided reaction sites for organic-inorganic bonding resulting in homogeneous and transparent hybrid films (Pan et al., 2018).

In medicine, L-2HG is used as a diagnostic biomarker for the characterization of various cancer types (Shim et al., 2014). Under oxygen limitation, L-2HG accumulates as metabolic intermediate in healthy as well as in malignant animal cells (Shim et al., 2014; Oldham et al., 2015; Shelar et al., 2018) facilitating a physiological adaption to hypoxic stress. The accumulation of L-2HG is triggered by increasing concentrations of 2-oxoglutarate (2-OG), which is caused by tricarboxylic acid cycle dysfunction and increased mitochondrial reducing potential. As consequence, the increased cellular L-2HG concentrations lead to inhibition of the electron transport and glycolysis compensating the effects of mitochondrial reductive stress induced by hypoxia (Oldham et al., 2015). The stereospecific reduction of 2-oxoglutarate to L-2HG is catalyzed by lactate dehydrogenase (LDH) and malate dehydrogenase (MDH) under hypoxic stress conditions (Intlekofer et al., 2017). In contrast, the formation of its enantiomer D-2-hydroxyglutarate (D-2HG) is catalyzed by a mutated version of isocitrate dehydrogenase 1 or 2 (IDH1/2) contributing to the pathogenesis of cancer, whereas L-2HG biosynthesis does neither involve IDH1 nor IDH2. Albeit both enantiomers of 2-hydroxyglutarate display an inhibitory effect on 2-oxoglutarate-dependent enzymes involved in diverse biologic processes (Chowdhury et al., 2011; Xu et al., 2011), their stereospecific biosynthesis differs. Thus, bio-based routes enable stereospecific synthesis of either D-2HG or L-2HG, which is preferred over chemically synthesized racemic D,L-2HG. Beyond the occurrence as a product of cellular redox stress, L-2HG also plays a role in plants and eukaryotic cells as part of the mitochondrial metabolic repair mechanism (Hüdig et al., 2015). A side reaction of MDH yields small concentrations of L-2HG by reduction of 2-oxoglutarate. The mitochondrial FAD-containing oxidase L-2-hydroxyglutarate dehydrogenase (L2HGDH, EC 1.1.5.13) oxidizes L-2HG and the electrons produced in the reaction are transferred to the mitochondrial electron transport chain through the electron transfer protein (ETF) (Wanders et al., 1997; Rzem et al., 2004). An enzyme homologous to L2HGDH was also described in the model organism *Escherichia coli* (Knorr et al., 2018).

In bacteria, L-2HG mainly accumulates under carbon starvation conditions (Marschall et al., 1998; Metzner et al., 2004; Knorr et al., 2018). It is also an intermediate of the L-lysine degradation pathway in Pseudomonadaceae (Zhang et al., 2018; Thompson et al., 2019). Glutarate is the direct precursor of L-2HG. Hydroxylation of glutarate to L-2HG is catalyzed by a highly regio- and stereospecific Fe(II)/2oxoglutarate-dependent dioxygenase CsiD (EC 1.14.11.64, also named glutarate hydroxylase) (Hibi and Ogawa, 2014). The coproduct succinate is metabolized via the TCA-cycle. Since L-2-hydroxyglutarate oxidase (LghO) regenerates 2-oxoglutarate by oxidation of L-2HG (Knorr et al., 2018; Herr et al., 2019), the combined activities of CsiD and LghO convert the C5dicarboxylic acid glutarate to the C4-dicarboxylic acid succinate. Thus, the absence of LghO is required for overproduction of L-2HG from glutarate.

Glutarate can be derived from L-lysine by four pathways that converge to 5AVA, which is converted to glutarate by GABA/5AVA aminotransferase (GabT) and the succinate/glutarate semialdehyde dehydrogenase (GabD). The first pathway from L-lysine to 5AVA employs L-lysine-αoxidase (RaiP) from Scomber japonicus that catalyzes oxidative desamination of L-lysine using molecular oxygen followed by spontaneous decarboxylation (Cheng et al., 2020). The second pathway to 5AVA combines oxidative decarboxylation by L-lysine monooxygenase using molecular oxygen followed by desamidation by γ -aminovaleramidase from *P. putida* (Adkins et al., 2013). The third pathway is based on L-lysine decarboxylase from E. coli, putrescine oxidase PuO from Rhodococcus qingshengii and the y-aminobutyraldehyde dehydrogenase from E. coli that catalyze decarboxylation, oxidative deamination using molecular oxygen and NADdependent oxidation (Haupka et al., 2020). The fourth pathway does not require molecular oxygen as it cascades L-lysine decarboxylase, 2-oxoglurate-dependent putrescine/cadaverine transaminase PatA, and NAD-dependent γ- aminobutyraldehyde dehydrogenase PatD from E. coli (Jorge et al., 2017). The pathway combinations for LdcC-PuO-PatD-GabT-GabD and LdcC-PatA-PatD-GabT-GabD couple conversion of lysine to glutarate in one or two transaminase reactions, respectively, that generate glutamate from 2-oxoglutarate. This coupling enabled flux enforcement, i.e., growth requires production of glutarate, which was achieved by deletion of gdh, the gene for the major ammonium assimilating enzyme L-glutamic acid dehydrogenase (Pérez-García et al., 2018; Haupka et al., 2020).

Since biosynthesis of L-2HG from glutarate requires molecular oxygen, glutarate biosynthesis from L-lysine by a pathway independent of molecular oxygen was extended by glutarate hydroxylase. The pathways RaiP-GabT-GabD, DavA-DavB-GabT-GabD, and LdcC-PuO-PatD-GabT-GabD involve oxygenases (RaiP, DavA, and PuO, respectively), whereas the pathway LdcC-PatA-PatD-GabT-GabD does not contain an enzyme using molecular oxygen as substrate (Jorge et al., 2017). To construct the six-step cascade LdcC-PatA-PatD-GabT-GabD-CsiD (**Figure 1**), CsiD enzymes from *E. coli* MG1655,



(EC 1.14.1.64) CsiD enzymes (violet box) were sourced from either E. coli (HEGIuA), P. putida (HPGIuA), or Halobacillus sp. (HBGIuA).

P. putida KT2440 and *Halobacillus sp.* BA-2008 were tested since inspection of the genome sequence of *C. glutamicum* did not indicate that a CsiD homolog is encoded. The absence of LhgO from the *C. glutamicum* genome was considered beneficial since it indicated that L-2HG may not be degraded. This and the fact that we previously established efficient glutarate production employing the LdcC-PatA-PatD-GabT-GabD pathway provided an ideal starting point to establish fermentative production of L-2HG from renewable resources. This concept may be transferred to fermentative production of the stereoisomer D-2HG by extending the glutarate pathway with enzymes for conversion of glutarate to D-2HG.

MATERIALS AND METHODS

Microorganisms and Cultivation Conditions

E. coli DH5 α strain was used as a host for cloning (Hanahan, 1985), S17-1 for transconjugation (Schäfer et al., 1994). Both strains were grown in lysogeny broth (LB) at 37 °C and supplemented with antibiotics (50 µg mL⁻¹ kanamycin, 100 µg mL⁻¹ spectinomycin, 10 µg mL⁻¹ tetracycline) when appropriate. *C. glutamicum* ATCC 13032 derived strains were cultivated in brain heart infusion with 0.5 M sorbitol (BHIS), or CGXII minimal medium (Eggeling and Bott, 2004) supplemented with 1 mM IPTG when appropriate. All bacterial strains and plasmids are listed in **Tables 1**, **2**. For standard growth experiments in CGXII medium with *C. glutamicum*, overnight cultures in 10 mL BHIS were harvested and washed

twice in TN-buffer (50 mM Tris-HCl, 50 mM NaCl, pH 6.3) before inoculation to an OD_{600} of 1 and supplementation with 40 g L⁻¹ glucose as a sole carbon source. The cultivations in the BioLector microfermentation system (m2p-labs, Baesweiler, Germany) were performed in 3.2 mL FlowerPlates at 1,100 rpm with a filling volume of 1,000 μ L.

To test for the utilization of L-2HG as carbon source, 12 mM L-2HG and 10 mM glucose were added, respectively, and the wildtype was inoculated to an initial OD₆₀₀ of 0.5. For optimization of the iron concentration in CGXII minimal medium the respective concentrations of iron-(II)-sulfate (0–3 mM) were supplemented. To test the inhibitory effect of glutarate on L-2HG production, 20 or 40 mM of glutarate (pH 7.0) was added to CGXII medium, respectively.

For L-2HG production from wheat sidestream concentrate (WSC; obtained from Jäckering, Hamm, Germany) overnight cultures in 10 mL BHIS were harvested and washed twice in TN-buffer (50 mM Tris-HCl, 50 mM NaCl, pH 6.3) before inoculation to an OD₆₀₀ of 1. The medium consisted of 246 g L⁻¹ WSC, 20 g L⁻¹ ammonium sulfate as nitrogen source, 42 g L⁻¹ MOPS as buffer and 2 mM iron-(II)-sulfate (added as CsiD is an iron-containing enzyme). Growth in 10 mL Duetz microcultivation plates (Kuhner Shaker GmbH, Herzogenrath, Germany) was performed with different sandwich covers and culture volumes to alter the oxygen supply. Under "low oxygen" cultivation, cells were grown in 3 mL at 220 rpm in an Ecotron ET25-TA-RC (Infors HT, Einsbach, Germany) and plate sandwich covers for low evaporation (1.2 mm hole diameter) were used. For "high oxygen" supply the culture

TABLE 1 | Bacterial strains used in this study.

Strain	Relevant characteristics	References	
E. coli			
DH5α	Δ lacU169 (ϕ 80lacZ Δ M15), supE44, hsdR17, recA1, endA1, gyrA96, thi-1, relA1	Hanahan, 1985	
S17-1	recA, pro, hsdR, RP4- 2Tc::Mu Km::Tn7 integrated into the chromosome	Simon et al., 1983	
C. glutamicum			
WT	C. glutamicum ATCC13032	ATCC	
GRLys1 (DM1933∆CGP123)	C. glutamicum ATCC13032 with modifications: Δpck, pyc ^{P458S} , hom ^{V59A} , 2 copies of lysC ^{T311I} , 2 copies of asd, 2 copies of dapA, 2 copies of dapB, 2 copies of ddh, 2 copies of lysA, 2 copies of lysE, in-frame deletion of prophages CGP1 (cg1507-cg1524), CGP2 (cg1746-cg1752) and CGP3 (cg1890-cg2071)	Unthan et al., 2015	
GSLA2G	In-frame deletions of sugR, IdhA, snaA and cgmA, gdh in GRLys1	Pérez-García et al., 2018	
GluA	GSLA2G(pVWEx1-ldcC) (pEKEx3-patDA) (pEC-XT99A-gabTD [*])	Pérez-García et al., 2018	
GluA∆ <i>sucE</i>	GSLA2G <i>ΔsucE</i> (pVWEx1 <i>-ldcC</i>) (pEKEx3 <i>-patDA</i>) (pEC-XT99A- <i>gabTD</i> *)	This study	
GluA2	GSLA2GΔP _{suce} ::P _{tuf} (pVWEx1- <i>ldcC</i>)(pEKEx3- <i>patDA</i>)(pEC-XT99A- <i>gabTD</i> *)	This study	
HEGluA	GSLA2G(pVWEx1- <i>ldcC-csiD^{Ec}</i>) (pEKEx3- <i>patDA</i>) (pEC-XT99A- <i>gabTD</i> [*])	This study	
HBGluA	GSLA2G (pVWEx1-ldcC-csiD ^{Hb}) (pEKEx3-patDA) (pEC-XT99A-gabTD [*])	This study	
HPGluA	GSLA2G (pVWEx1-IdcC-csiD ^{ep})(pEKEx3-patDA)(pEC-XT99A-gabTD*)	This study	
HPGluA∆ <i>sucE</i>	GSLA2G∆ <i>sucE</i> (pVWEx1 <i>-ldcC-csiD^cp</i>) (pEKEx3- <i>patDA</i>) (pEC-XT99A- <i>gabTD</i> *)	This study	
HPGluA2	IUA2 GSLA2GΔP _{sucF} ::P _{tuf} (pVWEx1- <i>IdcC-csiD^{Pp}</i>) (pEKEx3- <i>patDA</i>) (pEC-XT99A- <i>gabTD</i> *)		

TABLE 2 | Plasmids used in this study.

Plasmid	Relevant characteristics	References	
oK19mobsacB	Kan ^R , mobilizable <i>E. coli</i> vector mutagenesis (<i>orlV, sacB</i>)		
		Schäfer et al., 1994	
oK19mobsacB- Δ sucE	pK19mobsacB with a deletion construct sucE	This study	
$bK19mobsacB-\Delta P_{sucE}::P_{tuf}$	pK19 <i>mobsacB</i> with a promoter exchange of <i>sucE</i> with the strong <i>tuf</i> -promoter and an optimized RBS	This study	
oWWEx1	Kan ^R , <i>C. glutamicum/E. coli</i> shuttle vector (P _{tac} , <i>lacl</i> ^q)	Peters-Wendisch et al., 2001	
oVWEx1-IdcC	pVWEx1 expressing <i>IdcC</i> from <i>E.coli</i> MG1655	Pérez-García et al., 2018	
oVWEx1 <i>-ldcC-csiD^{Ec}</i>	pVWEx1 expressing IdcC and csiD from E.coli MG1655	This study	
oVWEx1 <i>-ldcC-csiD^{Hb}</i>	pVWEx1 expressing <i>ldcC</i> from <i>E.coli</i> MG1655 and <i>csiD</i> from <i>Halobacillus sp.</i> BA-2008	This study	
oVWEx1 <i>-ldcC-csiD^{Pp}</i>	pVWEx1 expressing <i>ldcC</i> from <i>E.coli</i> MG1655 and <i>csiD</i> from <i>Pseudomonas putida</i> KT2440	This study	
DEC-XT99A	Tet ^R , <i>C. glutamicum/E. coli</i> shuttle vector (P_{trc} , <i>lacl</i> ^q , pGA1, ori V_{Cg})	Kirchner and Tauch, 2003	
bEC-XT99A- <i>gabTD</i> *	pEC-XT99A expressing <i>gabT</i> and <i>gabD</i> ^{P134L} from <i>P. stutzerii</i> ATCC17588	This study	
DEKEX3	${ m Spec}^{ m R},$ C. glutamicum/E. coli shuttle vector (${ m P}_{tac}$ lacl ^q pBL1, oriV _{Ec})	Stansen et al., 2005	
EKEx3- <i>patDA</i> pEKEx3, expressing <i>patD</i> and <i>patA</i> from <i>E. coli</i> MG1655		Pérez-García et al., 2018	

Km^R: kanamycin resistance. Spec^R: spectomycin resistance. Tet^R: tetracycline resistance.

Primer	Sequence (5'-3')	Description		
ldcC-fw	CCTGCAGGTCGACTCTAGAGG ATTCCGAAAGGAGGCCCTTCAG ATGAACATCATTGCCATTATGGG	Construction of pVWEx1 <i>-ldcC-</i> <i>csiD</i>		
ldcC-csiD ^{Ec} -rv	CCTTTTGTATTCTTGTATTGGCG TTTTTATCCCGCCATTTTTAGG			
csiD ^{Ec} -fw	CCTAAAAATGGCGGGATAA AAACGCCAATACAAGAAT ACAAAAGGAGGTAATTTT ATGAATGCACTGACCGCCG	Construction of pVWEx1- <i>ldcC</i> - <i>csiD</i> ^{Ec}		
csiD ^{Ec} -rv	GAATTCGAGCTCGGTACCCGGGGAT CTTACTGATGCGTCTGGTAGT			
ldcC-csiD ^{Pp} -rv	GTCCTGTTAACAGGACTAATTAT AATTATCCCGCCATTTTTAGG			
csiD ^{Pp} -fw	CCTAAAAATGGCGGGATAA <i>TTATAATTAGTCCTGTTAACAG GACATCAAAGGAGGTTTTTTT</i> ATGAACGCCTTTACGCAG	Construction of pVWEx1 <i>-ldcC-</i> <i>csiD^{Pp}</i>		
csiD ^{Pp} -rv	GAATTCGAGCTCGGTACCCGGGGATC TTATTGACCGCGCTGGTAC			
ldcC-csiD ^{Hb} -rv	<i>TTCCTTTAAGTTATACTTTCGTTAA</i> TTATCCCGCCATTTTTAGG			
csiD ^{Hb} -fw	CCTAAAAATGGCGGGATAA <i>TTAACGAAAGTATA ACTTAAAGGAACCACGTATTT</i> ATGTGCGCAGTAGAAATG	Construction of pVWEx1- <i>IdcC-</i> <i>csiD^{Hb}</i>		
csiD ^{Hb} -rv	GAATTCGAGCTCGGTACCCGGGGATC CTATTGAAGGAATCGTCC			
ldcC-seq1	TGAACGATGTAGTGCCAGTC			
ldcC-seq2	GCAATGGGATTATTGCGTGG	Sequencing of pVWEx1 <i>-ldcC-</i> <i>csiD</i>		
ldcC-seq3	CAGGCAGAATCGAAGTGA			
csiD ^{Ec} -seq	CCGATTACGTGCTGATG	Sequencing of pVWEx1 <i>-ldcC-</i> <i>csiD^{Ec}</i>		
csiD ^{Pp} -seq	GATCTGGTTCACGAAC	Sequencing of pVWEx1 <i>-ldcC-</i> <i>csiD^{Pp}</i>		
csiD ^{Hb} -seq	GTAGGTTCCATCAGTG	Sequencing of pVWEx1 <i>-ldcC-</i> <i>csiD^{Hb}</i>		
PSucEA	<i>GCATGCCTGCAGGTCGACTCTAGA</i> GGCGTGACGTGTACAAGCGCG			
PSucEB	TACGCGCCTACTGACACGCTAAAA CTTAAGCCTCGCCCTTGCGTTC	Construction of pK19 <i>mobsacB_</i>		
P _{tuf} -fw	GGCTGAACGCAAGGGCGAGGCTTAA GTTTTAGCGTGTCAGTAGGC			
P _{tuf} -rv	AAGGAAGCTCAT AAAAATACCTCCCCCAGTGTTCGTG CCGTCGCCCCGGCGACGAGTTTA GTTACTGAATCCTAAGGGCA			
PSucEC				
PSucED	AATTCGAGCTCGGTACCCGGGGATC GAATAACGATGAGCACACCG			

(Continued)

TABLE 3 | Continued

Primer	Sequence (5'-3')	Description		
PSucEE	GACTCGCTCACAAATGTGG	Verification of Promoter exchange ΔP_{sucE} ::P _{tuf}		
PSucEF	GAATTGCTCACCGTCTCG			
SucEA	GCATGCCTGCAGGTCGACTCTAGAG GTGGCACCTGGTGTTCCAG			
SucEB	TTTGGGCGGCCAGGATCTTTGCGAT TTCTACAAGGAAGCTCAC	Construction of pK19 <i>mobsacB_</i> ∆ <i>sucE</i>		
SucEC	GAATGGGTGAGCTTCCTTGTAGAA ATCGCAAAGATCCTGGC			
SucED	AATTCGAGCTCGGTACCCGGGGATC CGAATGGATTGGTCAGGG			
SucEE	CTGCTGGTTGGGCTGTGG	Verification sucE deletion		
SucEF	GTTAATCATGAGGCGTCG	Verification sucE deletion		

Overlaps to the vector are underlined, the ribosome binding site is indicated in italic and amino acid exchanges are shown in bold.

volume was decreased to 2 mL and standard plate sandwich covers (2.5 mm hole diameter) were used. Growth was monitored by determination of the OD₆₀₀ with a V-1200 Spectrophotometer (VWR, Radnor, PA, USA).

Molecular Biology Methods

Genomic DNA of C. glutamicum, E. coli and P. putida were isolated as described previously (Eikmanns et al., 1994). The gene csiD from Halobacillus sp. BA-2008 was codonharmonized (Haupka, 2020) and synthesized with an optimized ribosomal binding site by Synbio Technologies (South Brunswick Township, New Jersey, United States of America). The standard molecular methods including plasmid isolation, molecular cloning and transformation of E. coli by heat shock and of C. glutamicum by electroporation with plasmid DNA were performed as described before (Eggeling and Bott, 2004). DNA sequences were amplified with the ALLin HiFi DNA Polymerase (HighQu, Kraichtal, Germany) using plasmid or genomic DNA as template. The oligonucleotides used in this study are listed in Table 3. The gene *ldcC* was amplified from the vector pVWEx1-ldcC (Pérez-García et al., 2018), whereas the different csiD genes were amplified from genomic DNA of the respective organisms and assembled together into BamHI-digested pVWEx1 by Gibson Assembly, using the respective primers. The constructed plasmids were transferred into C. glutamicum by transformation. For deletion, plasmid pK19mobsacB (Schäfer et al., 1994), digested with BamHI was assembled with amplified DNA fragments flanking the gene sucE (cg2425) using Gibson Assembly and was transferred into E. coli S17-1 to follow a protocol for gene deletion routinely applied (Eggeling and Bott, 2004). For replacement of the native promoter by the stronger *tuf*-promoter, the *tuf*-promoter and the flanking regions of the promoter of sucE were amplified from

genomic DNA of *C. glutamicum* and assembled with the digested plasmid pK19*mobsacB*. For higher expression rates the start codon of *sucE* was changed from GTG to ATG and an optimized ribosome binding site (RBS) was included (Salis, 2011).

Coupled in vitro Activity of GabT and GabD

The apparent activities of GABA transaminase GabT and succinate semialdehyde oxidoreductase GabD were assayed in combination by monitoring NADPH formation after the addition of 2-oxoglutarate. The preparation of the crude extract was carried out as previously described (Pérez-García et al., 2018). The 1 mL assay mix contained 150 mM phosphate buffer (pH 9.0), L-2HG (2, 4, 8, 12 mM) or NaCl as control, 0.1 mM pyridoxal-5'-phosphate, 1 mM NADP⁺, 20 mM 5AVA, and 0.5 mg mL⁻¹ crude extract. The reaction was started by the addition of 15 mM 2-oxoglutatarate. Protein concentrations were determined with the Bradford assay kit (Bio-Rad Laboratories, Hercules, CA, United States) using BSA (bovine serum albumin) as standard. The formation of NADPH was monitored photometrically at 340 nm and 30°C for 3 min using a Shimadzu UV-1202 spectrophotometer (Shimadzu, Duisburg, Germany).

Quantification of Amino Acids, Diamines, and Carboxylic Acids

The quantification of extracellular amino acids and their derivatives, carbohydrates and carboxylic acids in the cultivation medium was performed with a high-performance liquid chromatography system (1200 series, Agilent Technologies Deutschland GmbH, Böblingen, Germany). After centrifugation of 1 mL of cell cultures at 14,000 rpm for 10 min the supernatant was stored at -20° C prior to analysis. Analysis of L-lysine, 5AVA and the diamine cadaverine was performed by an automatic pre-column derivatization with ortho-phthaldialdehyde (OPA) and separated on a reversed phase HPLC using pre- and main column (LiChrospher 100 RP8 EC-5 μ , 125 \times 4.6 mm, CS Chromatographie Service GmbH) with L-asparagine as internal standard (Schneider and Wendisch, 2010). Detection of the fluorescent derivatives was carried out with a fluorescence detector with an excitation wavelength of 230 nm and an emission wavelength of 450 nm. Glutarate and L-2HG and glucose concentrations were measured with an amino exchange column (Aminex, 300×8 mm, $10 \,\mu$ m particle size, 25 Å pore diameter, CS Chromatographie Service GmbH) under isocratic conditions as described previously with a flow of 0.8 mL min⁻¹ (Schneider et al., 2011). The substances were detected with a refractive index detector (RID G1362A, 1200 series, Agilent Technologies) and a diode array detector (DAD G1315B, 1200 series, Agilent Technologies) at 210 nm.

Fermentative Production

A baffled bioreactor with total a volume of 3.6 L was used (KLF, Bioengineering AG, Switzerland). Three six-bladed Rushton turbines were placed on the stirrer axis with a distance from the bottom of the reactor of 6, 12, and 18 cm. The aspect ratio of the reactor was 2.6:1.0 and the stirrer to reactor diameter ratio was 0.39. Automatic control of the stirrer speed between 400 and 1500 rpm kept the relative dissolved oxygen saturation at 30%.

A constant airflow of 1 or 2 NL min^{-1} was maintained from the bottom through a sparger, corresponding to an aeration of 0.5 and 1 vvm, respectively.

The pH was kept constant at 7.0 \pm 0.1 by automatic addition of phosphoric acid [10% (v/v)] and potassium hydroxide (4 M). The temperature was maintained at 30°C. To prevent foaming $0.6 \text{ mL} \text{ L}^{-1}$ of the antifoam agent AF204 (Sigma Aldrich, Darmstadt, Germany) was added and a mechanical foam breaker was present to serve as an additional foam control. The fermentation was performed with a head space overpressure of 0.2 bar. The initial working volume of 2 L was inoculated to an OD₆₀₀ of 1.5 from a shake flask pre-culture in CGXII minimal medium supplemented with 40 g L^{-1} glucose, 1 mM IPTG and 2 mM FeSO₄. Samples were collected by an autosampler and cooled down to 4°C until further use. The feed consisted only of 600 g L⁻¹ glucose ($\rho = 1.21$ kg m⁻³) and was started 4 h after the cells reached the late stationary phase, indicated by the pO_2 rising above 80% again after the initial growth phase. The glucose feed was applied for 5 min with a flow of 1.2 mL min^{-1} when the pO₂ surpassed 60%. Further feed was only added, when the pO₂ decreased to 30% after the addition of the feed solution to prevent oversaturation with glucose.

RESULTS

Establishing *de novo* Biosynthesis of L-2HG by *C. glutamicum*

As *C. glutamicum* can utilize some organic acids (Wendisch et al., 2016), the response of *C. glutamicum* WT to L-2HG was determined. When L-2HG was added to mineral salts medium as sole carbon source instead of glucose, no growth of *C. glutamicum* was observed. Moreover, when L-2HG plus glucose were assayed, *C. glutamicum* WT utilized glucose, but did not degrade L-2HG as revealed by HPLC analysis of the culture supernatants (data not shown). Thus, L-2HG does neither serve as sole nor as combined carbon source for *C. glutamicum*, which can be considered as a suitable host for production of L-2HG.

L-2HG is not a known metabolite of C. glutamicum, but it has been engineered for overproduction of glutarate (Pérez-García et al., 2018). Inspection of the genome of C. glutamicum did not indicate the presence of a gene coding for an enzyme hydroxylating glutarate. In some bacteria like E. coli (Marschall et al., 1998; Knorr et al., 2018; Herr et al., 2019), P. putida, and Halobacillus sp. (Thompson et al., 2019), glutarate hydroxylase, also known as glutarate dioxygenase (EC 1.14.11.64), uses molecular oxygen to hydroxylate glutarate to L-2HG with concomitant decarboxylation of 2-oxoglutarate to succinate. Thus, the respective csiD genes from E. coli MG1655, P. putida KT2440 and Halobacillus sp. BA-2008 were overexpressed in synthetic operons with *ldcC* from *E. coli*. The resulting pVWEx1plasmids were used to transform C. glutamicum GSLA2G (pEKEx3_patDA) (pEC-XT99A_gabTD*). The resulting strains were named HEGluA, HPGluA, and HBGluA, respectively. In production experiments, the three strains produced and secreted L-2HG. After 96 h C. glutamicum strain HEGluA accumulated 14 \pm 0 mM L-2HG, strain HBGluA 7 \pm 1 mM L-2HG, and strain HPGluA 22 \pm 2 mM L- 2HG (**Figure 2**). The strains showed comparable maximal growth rates of 0.13 \pm 0.00 h^{-1}, but strain HPGluA showed a lag-phase of 24 h. Taken together, a proof-of-principle for production of L-2HG by *C. glutamicum* was demonstrated.

Role of the Succinate Exporter SucE in Export of Glutarate and L-2HG

SucE is known to play a role in the export of the dicarboxylic acid succinate out of the *C. glutamicum* cell (Huhn et al.,

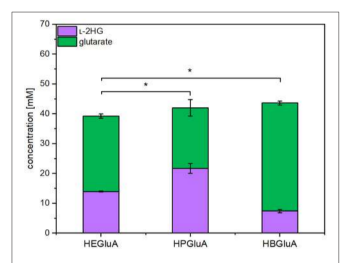


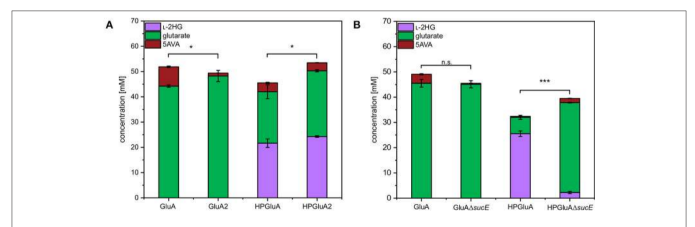
FIGURE 2 | Product titers of L-2HG and its precursor glutarate obtained with strains overproducing different glutarate hydroxylases. *C. glutamicum* strains HEGluA, HPGluA, and HBGluA expressing *csiD* from *E. coli* MG1655, *P. putida* KT2440, and *Halobacillus sp.* BA-2008, respectively, were grown in 40 g L⁻¹ glucose CGXII minimal medium supplemented with 1 mM IPTG in the microcultivation device BioLector. Values and error bars represent means and standard deviations from 3 replicate cultivations with supernatants analyzed after 96 h. Statistical significance was assessed by Student's paired *t*-testing (*p < 0.05, n.s, not significant).

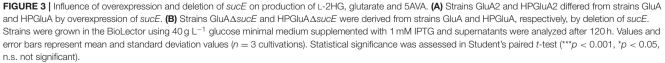
2011). Since L-2HG and glutarate are also dicarboxylic acids, the role of SucE in the export of L-2HG and glutarate was studied. To this end, the *sucE* gene was either overexpressed by replacing the native promoter with the stronger *tuf*-promoter or deleted. The glutarate producing strain GluA2 that overexpressed *sucE* produced significantly more glutarate than the control strain GluA (44 \pm 0 vs. 48 \pm 2 mM; **Figure 3A**). Similarly, upon overexpression of *sucE*, production of L-2HG was slightly elevated (24 \pm 0 mM for HPGluA2 vs. 22 \pm 2 mM for HPGluA; **Figure 3A**) as well as production of the by-product glutarate (26 \pm 0 mM for HPGluA2 vs. 20 \pm 3 mM for HPGluA; **Figure 3A**). Thus, overexpression of *sucE* was beneficial for production of glutarate as well as L-2HG.

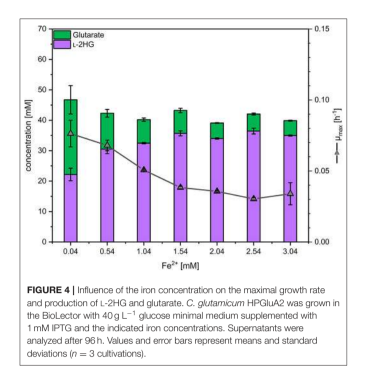
Deletion of *sucE* in glutarate producer GluA did not negatively affect glutarate production, however, growth was slowed (0.12 \pm 0.00 h⁻¹ for GluA vs. 0.07 \pm 0.00 h⁻¹ for GluA Δ *sucE*; **Figure 3B**). Thus, SucE is not the main export system for glutarate and at least one other export system is able to compensate for the lack of SucE regarding glutarate export. Importantly, deletion of *sucE* reduced production of L-2HG more than 10-fold (2 \pm 0 mM for HPGluA Δ *sucE* vs. 26 \pm 1 mM for HPGluA; **Figure 3B**), while the growth rate was not significantly impacted (0.09 \pm 0.00 h⁻¹ for HPGluA vs. 0.08 \pm 0.01 h⁻¹ for HPGluA Δ *sucE*). The findings that *sucE* overexpression positively affected L-2HG production and that deletion of *sucE* dramatically reduced L-2HG production suggested that SucE may be active as export system for L-2HG.

Enhanced Conversion of Glutarate to L-2HG by Increased Iron Concentrations

As the glutarate hydroxylase CsiD is an Fe²⁺-dependent metalloenzyme (Herr et al., 2019; Thompson et al., 2019), the concentration of iron (II) in the media was varied and the effect on production of L-2HG determined. Concentrations of 0.5 to 3 mM FeSO₄ were added on top of the standard concentration (0.037 mM) in CGXII minimal medium







(Eggeling and Bott, 2004). Three trends were observed: with increasing iron concentrations the growth rate and the formation of glutarate as by-product were reduced, while production of L-2HG was significantly increased (**Figure 4**). At an iron concentration of 2.04 mM, 5 ± 0 mM glutarate and 34 ± 0 mM L-2HG accumulated in the supernatant (**Figure 4**), thus, about 87% of glutarate were converted to L-2HG, while at the standard iron concentration L-2HG and glutarate accumulated to about equimolar concentrations. The growth inhibitory effect of elevated iron concentrations may be due to increased production of L-2HG and/or other effects elicited by higher iron concentrations. Thus, 2.04 mM was choosen as the optimal iron concentration for further experiments to find the best compromise between L-2HG titer and volumetric productivity.

Inhibitory Effects on Key Enzymes in L-2HG Pathway

The finding that at higher iron concentrations less glutarate and more L-2HG accumulated, while the combined concentration of glutarate plus L-2HG was highest at the lowest iron concentration indicated bottlenecks. These may arise due to inhibition of glutarate hydroxylase CsiD by its product L-2HG (Knorr et al., 2018) and/or due to inhibition of enzymes of glutarate biosynthesis. The transaminases PatA from *E. coli* and GabT from *P. stutzerii* are crucial for glutarate biosynthesis (Pérez-García et al., 2018) and some transaminases are known to be inhibited by L-2HG, which blocks binding of the substrate 2oxoglutarate (McBrayer et al., 2018). To approach this problem, enzyme activity assays were performed with crude extracts of *C. glutamicum* GluA. A combined activity assay of GABA transaminase GabT and succinate semialdehyde dehydrogenase

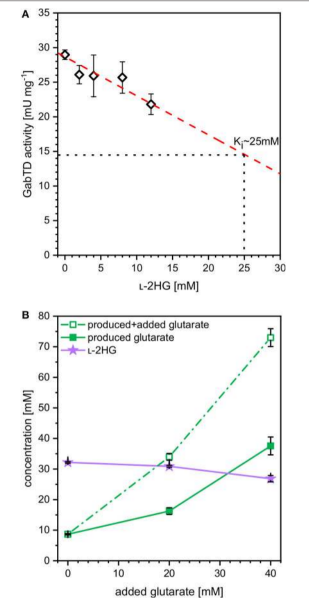


FIGURE 5 | Influence of L-2HG on the combined *in vitro* enzyme activities of GABA transaminase GabT and succinate semialdehyde dehydrogenase GabD
(A) and influence of extracellularly added glutarate on production of L-2HG (B).
(A) Crude extracts of GluA were assayed for combined *in vitro* enzyme activities of GABA transaminase GabT and succinate semialdehyde dehydrogenase GabD in the presence of increasing concentrations of L-2HG.
(B) Strain HPGluA2 was cultivated in the BioLector with 40 g L⁻¹ glucose minimal medium supplemented with 1 mM IPTG, 2 mM iron (II)-sulfate and increasing concentrations of glutarate (0, 20, 40 mM). Supernatant concentrations of L-2HG (filled violet triangles), glutarate (open green squares) as well as the net glutarate concentrations produced in addition to the added glutarate concentration (closed green squares) were determined after 96 h and are given as means and standard deviations of three independent cultivations.

GabD was performed as described previously (Pérez-García et al., 2018). In the presence of increasing concentrations of L-2HG (0–12 mM) the combined GabT and GabD activity was

reduced. By extrapolation, it was determined that 25 mM L-2HG resulted in half-maximal GabTD activity (**Figure 5A**). Thus, L-2HG negatively affects production of its precursor glutarate.

In order to determine if increased glutarate concentrations are beneficial for production of L-2HG, production of L-2HG was determined in the absence or presence of extracellularly added glutarate (0, 20, or 40 mM). Notably, externally added glutarate slightly reduced production of L-2HG, whereas glutarate accumulation was increased (**Figure 5B**). When 0, 20, or 40 mM glutarate were added to the medium before inoculation, additional glutarate was produced: 9, 16, and 38 mM, respectively (**Figure 5B**). Thus, extracellular glutarate addition exerts a positive effect on glutarate production. The slight reduction of L-2HG production upon addition of extracellular glutarate may be due to substrate inhibition of glutarate hydroxylase CsiD.

L-2HG Production at Bioreactor Scale

In order to test if L-2HG production by strain HPGluA2 is stable at larger volumes, 2 L scale bioreactor fermentations were performed. Two aeration rates were tested: 0.5 and 1 vvm (**Figure 6**). Under both conditions cells grew with similar growth rates (0.5 vvm: 0.07 h⁻¹, 1 vvm: 0.08 h⁻¹) and displayed a similar entry into the stationary phase after 56 h. Oscillations in rDOS was due to overregulation of the stirrer, an often observed disadvantage of PID controllers, especially, when combined with the rDOS probes that are highly sensitive to fluctuations. However, almost 2-fold more biomass was formed at the higher

aeration rate (0.5 vvm: 6.3 g L^{-1} , 1 vvm: 9.9 g L^{-1}). Accumulation of the precursor glutarate was growth-coupled and ceased upon entry into the stationary phase (titers of around 10 mM at 0.5 vvm and 13 mM at 1 vvm), whereas production of L-2HG was delayed and started after 30 h. While L-2HG production at an aeration rate of 0.5 vvm reached the highest titer of 16 mM at 120 h (**Figure 6A**), L-2HG production at an aeration rate of 1 vvm stopped at a titer of 6 mM shortly after glucose was fully depleted (**Figure 6B**). Thus, the latter fermentation was switched to feeding mode to supply the culture with more carbon source. As consequence, the strain grew to a 2-fold higher biomass concentration of 16.3 g L⁻¹ and produced fourfold more L-2HG reaching a titer of 24 mM (**Figure 6B**).

Comparing the different cultivations in batch mode (**Table 4**) it becomes obvious that the L-2HG product yield showed an inverse relationship with the aeration rate, as its production was highest in the not actively aerated microcultivation system. Although HPGluA2 grew 2-fold slower in the microcultivation system, the volumetric productivity was higher than in the bioreactors (**Table 4**). By contrast, growth proceeded to the highest biomass concentration and the highest biomass yield was observed when aeration was highest (**Table 4**).

L-2HG Production From Wheat Sidestream Concentrate

Sustainable processes based on renewable feedstock are sought after. Therefore, it was tested if L-2HG can be produced from

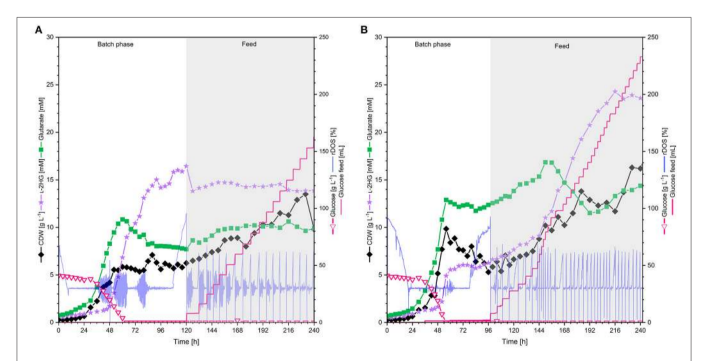
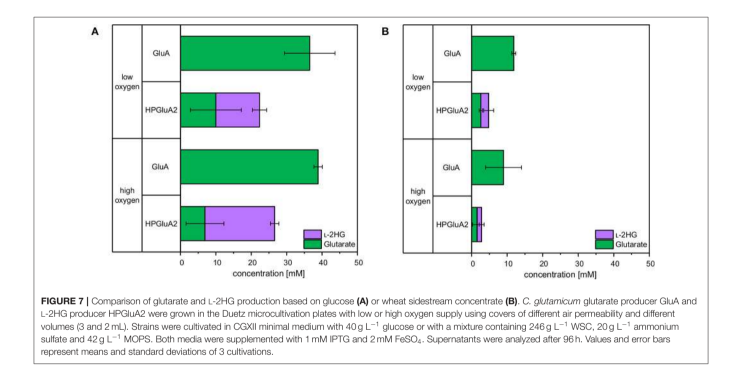


FIGURE 6 | L-2HG production by *C. glutamicum* HPGluA2 in fed-batch fermentation with (A) 0.5 vvm and (B) 1 vvm aeration rate. HPGluA2 was cultivated in CGXII minimal medium in fed-batch mode over 240 h, containing 40 g L⁻¹ glucose and feeding 600 g L⁻¹ glucose solution. L-2HG concentration is indicated in violet stars (mM), biomass concentration (CDW) is shown in black diamands (g L⁻¹), glucose concentration (g L⁻¹) is plotted as pink hollow triangles, and glutarate concentration (mM) in green squares, 600 g L⁻¹ glucose feed (mL) is plotted as pink line and the relative dissolved oxygen saturation (rDOS) is indicated in light blue (%). Cultivation was performed at 30°C and pH 7.0 regulated with 10% (*v/v*) H₃PO₄ and 4 M KOH. An overpressure of 0.2 bar was applied. 0.6 mL L⁻¹ of antifoam agent AF204 (Sigma Aldrich, Taufkirchen, Germany) was added to the medium manually before inoculation.

System	Mode/Phase	Aeration rate [vvm]	^μ max [h ⁻¹]	CDW [g L ⁻¹]	Titer [g L ^{−1}]	Y _{X/S} [g g ⁻¹]	Y _{P/S} [g g ⁻¹]	VP [g L ⁻¹ h ⁻¹]
Micro- Cultivation	Batch	_	0.04	4.3	5.0	0.11	0.13	0.05
Bioreactor	Batch	0.5	0.07	6.3	2.4	0.16	0.06	0.02
	Batch	1	0.08	9.9	0.9	0.25	0.02	0.01
	Fed-batch	1	0.07	16.3	3.5	0.09	0.02	0.01

TABLE 4 | Comparison of L-2HG process parameters during different cultivation strategies.

 μ_{max} , maximal growth rate; CDW, cell dry weight; $Y_{X/S}$, biomass yield; $Y_{P/S}$, product yield; VP, volumetric productivity.



a sidestream of industrial starch production (wheat sidestream concentrate; WSC). WSC contains hardly any starch, but various sugars like glucose, fructose, sucrose, raffinose, xylose and arabinose (D'Appolonia and Rayas-Duarte, 1994). Production of glutarate from WSC was also analyzed. To this end, C. glutamicum strains GluA and HPGluA2 were cultivated for 96 h either in CGXII medium with glucose or in WSC medium. Aeration was altered by different cultivation volumes and by using either "low oxygen" or "high oxygen" Duetz plates. For HPGluA2 both media were supplemented with 2 mM FeSO₄. Glucose-based glutarate production by strain GluA was comparable for both aeration schemes (37 \pm 7 and 39 \pm 1 mM; Figure 7), whereas it was lower in WSC medium (12 \pm 1 and 9 \pm 5 mM of glutarate under high and low oxygen supply, respectively; Figure 7). HPLC analysis of WSC using a refractive index detector did not allow to identify all peaks, but we could show that around 12 mM glucose and 6 mM maltose were utilized (data not shown) and 3 $\pm 0 \,\text{mM}$ glutarate and $2 \pm 1 \,\mathrm{mM}$ L-2HG accumulated under low oxygen conditions.

No significant differences with respect to L-2HG production by strain HPGluA2 with different oxygen supply were observed (**Figure 7**). Albeit leading to lower titers, production of L-2HG to concentrations around 2 mM were achieved. Thus, L-2HG production from the renewable feedstock wheat sidestream concentrate was demonstrated.

DISCUSSION

In this study, production of the sought after compound L-2HG by the industrial workhorse *C. glutamicum* was demonstrated after extension of lysine biosynthesis in a six-step cascade employing the metalloenzyme CsiD from *P. putida* as final step. The glucose-based process was stable in 2L bioreactor cultivations and a L-2HG titer of 3.5 g L^{-1} was obtained in fed-batch fermentation. Moreover, L-2HG production based on the renewable feedstock wheat sidestream concentrate was demonstrated.

L-2HG production was achieved by glutarate hydroxylase extending lysine biosynthesis in a six-step cascade, in which only the last committed step catalyzed by oxygen-dependent glutarate hydroxylase requires oxygen. Glutarate hydroxylases belong to the large family of non-heme Fe(II)- and 2oxoglutarate-dependendent oxygenases, which are essential for diverse biological functions. These enzymes form an Fe(IV)oxo intermediate to initiate oxidative transformations and can be assigned to four major types of reactions: hydroxylation, halogenation, ring formation and desaturation (Hausinger, 2004). The glutarate hydroxylases which have been studied in this research belong to the enzymes catalyzing a hydroxylation reaction at an unactivated carbon center by incorporation of molecular oxygen (Guengerich, 2015; Martinez and Hausinger, 2015). The CsiD enzymes from E. coli MG1655 and P. putida KT2440 have been recently characterized in different studies displaying their high specificities toward the native substrate glutarate with K_M-values around 0.65 mM in E. coli (Knorr et al., 2018) and around 0.15 mM for P. putida (Zhang et al., 2018). The K_M-values of around 0.1 mM for the co-substrate 2-oxoglutarate are comparable for both enzymes. The higher affinity of glutarate hydroxylase from P. putida for glutarate is reflected by the better conversion of glutarate to L-2HG observed in this study. Even though no K_M value for the glutarate hydroxylase from Halobacillus sp. BA-2008 has been determined vet, it was demonstrated that it converts 5 mM glutarate to L-2HG with comparable efficiency to the other tested hydroxylases with 2-oxoglutarate as co-substrate (Thompson et al., 2019). Although the capability of CsiD from Halobacillus sp. BA-2008 to produce L-2HG from glutarate was demonstrated in C. glutamicum, the efficiency of the codon harmonized version of CsiD was inferior compared to the enzymes derived from other organisms.

It could be demonstrated that L-2HG inhibits the combined GabTD activity with an inhibitory constant of about 25 mM (Figure 5A). Due to its structural similarity to 2-OG, L-2HG potentially inhibits transamination reactions (McBrayer et al., 2018) by competitive inhibition (Intlekofer et al., 2015). Here, possibly competitive inhibition of 2-OG dependent transaminase GabT by L-2HG may have limited L-2HG product titers. Potentially, this may be overcome by enzyme engineering of the 2-OG binding pocket of GabT for better differentiation of this substrate from L-2HG. We have chosen enzymes GabT and GabD from P. stutzeri as they performed better than those from C. glutamicum, P. putida and P. syringae regarding glutarate production (Pérez-García et al., 2018). Since we did not compare GabT and GabD enzymes from various sources for inhibition of their activities by L-2HG, identifying feedback resistant enzymes/variants of the GabT and GabD enzymes in future work may help to increase L-2HG production. On the other hand, glutarate hydroxylase CsiD is subject to weak product inhibition (Knorr et al., 2018). Therefore, CsiD from other sources or variants may be selected that exhibit no or reduced product inhibition in order to improve L-2HG production.

Moreover, a weak substrate inhibition on L-2HG production by glutarate could also be identified (**Figure 5B**). CsiD from *E. coli* was recently characterized and the reaction mechanism was described (Knorr et al., 2018; Herr et al., 2019). Since the substrate analogon *N*-oxalylglycine (NOG), a 2-OG mimic, inhibited CsiD (Knorr et al., 2018), this feedback inhibition may limit L-2HG production. However, since we did not assay feedback inhibition of various glutarate hydroxylases by L-2HG, future work to improve L-2HG production should involve the identification of feedback resistant CsiD enzymes/variants.

Surprisingly, extracellular addition of glutarate boosted glutarate production. This is unlikely due to effects on enzyme activities. *C. glutamicum* possesses chromosomal copies of *gabT* and *gabD* (Pérez-García et al., 2018; Haupka et al., 2020). The PucR-like regulator GabR that requires GABA as coactivator activates transcription of the *gabTDP* operon (Zhu et al., 2020). Possibly, glutarate mimics GABA as coactivator of GabR activating the endogenous *gabTDP* operon and increasing GabT and GabD enzyme levels in addition to plasmid-borne expression of *gabT* and *gabD* from *P. stutzeri*.

Cultivation schemes with higher oxygen supply reduced L-2HG production, but improved growth to high biomass concentrations. Glutarate hydroxylases are 2-OG-dependent enzymes and these metalloenzymes specifically require Fe²⁺ as cofactor for their function (Mitchell et al., 2017; Dunham et al., 2018). Previously, it has been shown that increased iron concentrations improve enzyme activity (Fukumori and Hausinger, 1993). Increased Fe²⁺ concentrations improved L-2HG product titers, but slowed growth significantly. This might be due to iron effects such as toxicity mainly due to the formation of hydroxyl radicals as oxidative stress response (Braun, 1997; Touati, 2000; Eid et al., 2017) and/or due to inhibition of 2-OG-dependent dioxygenases and methylases, which play an essential role in DNA/RNA repair (van den Born et al., 2009), by L-2HG (Low et al., 2001; Ledesma-García et al., 2016). Provision of oxygen and iron to the C. glutamicum cell are interlinked. Molecular oxygen is required for 2-OGdependent dioxygenases such as CsiD to form the Fe(IV)-oxo intermediate, but with too much oxygen decoupling occurs via "non-productive reactions" resulting in inactive Fe(III)containing enzymes (Hausinger, 2004) as observed with AlkB (Henshaw et al., 2004) and TfdA (Liu et al., 2001). Secondly, an "uncoupled turnover" of the co-substrate 2-OG may occur by decomposition of 2-OG to carbon dioxide and succinate. These "uncoupled reactions" might contribute to the lower titers of L-2HG obtained in 2L scale fermentative production with the highest oxygen supply. The addition of ascorbic acid is a promising option since ascorbate improved hydroxylase activity (Fukumori and Hausinger, 1993). Alternatively, production of L-2HG may be improved by concomitant overproduction of either glutathione (Liu et al., 2019) or ascorbate in C. glutamicum.

Our results suggested that export of L-2HG involves SucE (**Figure 3**). Surprisingly, glutarate export does not depend solely on SucE. Overexpression of *sucE* increased succinate production (Zhu et al., 2014) similar to increased production of glutarate and L-2HG observed here. Deletion of *sucE* revealed that SucE is the main export system for L-2HG, but not for glutarate. Production of L-2HG may benefit from re-uptake of glutarate secreted to the culture medium as by-product. However, the uptake system for glutarate is unknown. MctC is involved in the uptake of pyruvate,

propionate, and acetate, but it is not clear if it accepts glutarate (Jolkver et al., 2009).

Glutarate was observed as significant by-product. Unlike L-2HG, which likely is exported by SucE (s. above), glutarate is exported by YnfM (Fukui et al., 2019). Overexpression of *ynfM* improved production of glutarate, succinate, and 2-OG (Fukui et al., 2019; Han et al., 2020). Thus, deletion of *ynfM* is a suitable strategy to abolish export of glutarate as by-product of L-2HG production.

Alternative feedstocks are important to achieve sustainable biotechnological processes. In this respect, sidestreams are highly relevant. Glycerol is a sidestream of the biodiesel process and glycerol accumulates as stoichiometric by-product. C. glutamicum has been engineered to produce amino acids from pure and technical grade glycerol (Rittmann et al., 2008; Meiswinkel et al., 2013b). Access to nitrogenous sidestreams from the fishery industry such as glucosamine or N-acetylglucosamine is important if the target product is containing nitrogen atoms (Matano et al., 2014, 2016). For products lacking nitrogen atoms, access to lignocellulosics is pivotal (Gopinath et al., 2011; Buschke et al., 2013; Meiswinkel et al., 2013a; Hadiati et al., 2014; Matsuura et al., 2019). The focus of this study was a sidestream of industrial starch production. Although production was lower compared to glucose (30% for glutarate, 15% for L-2HG), media composition was easy and required only a buffer, a nitrogen source for growth and iron for glutarate hydroxylase. We envision improved production of L-2HG from wheat sidestream concentrate once access to sugars other than the native substrates glucose, fructose, maltose, and sucrose is achieved, e.g., to raffinose, xylose and arabinose, that are present in this feedstock (D'Appolonia and Rayas-Duarte, 1994). Strategies for access to the lignocellulosic pentoses xylose and arabinose have been established (Kawaguchi et al., 2008; Gopinath et al., 2011; Schneider et al., 2011; Meiswinkel et al., 2013a; Imao et al., 2017) and may prove useful for improved production of L-2HG from the alternative feedstock wheat sidestream concentrate. In this respect, provision of iron and/or iron chelators may prove essential as shown here for CsiD function.

The approach described here may be followed for production of the stereoisomer D-2HG. To this end, the glutarate pathway described here has to be extended with enzymes for conversion of glutarate to D-2HG. These enzymes may be sourced from lysine degrading bacteria as D-2HG occurs as intermediate in some of these pathways, e.g., in *E. coli* (Zhao and Winkler, 1996) and different *Pseudomonaceace* (Zhang et al., 2017; Guo et al., 2018; Thompson et al., 2019). Future work will reveal if this approach is suitable for fermentative production of D-2HG.

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DATA AVAILABILITY STATEMENT

The original contributions presented in the study are included in the article/supplementary material, further inquiries can be directed to the corresponding author/s.

AUTHOR'S NOTE

VFW wishes to dedicate this article in memoriam of scientific and personal friend Michael M. Goodin, Professor of Plant Pathology, University of Kentucky, Lexington, KY.

AUTHOR CONTRIBUTIONS

CP and VFW conceived and designed the experiments. CP constructed plasmids and strains, cultivated, and analyzed *C. glutamicum* strains and prepared a draft of the manuscript. CP and FM performed bioreactor experiments and evaluated data. AB conducted the growth experiment on wheat sidestream concentrate. CP, AB, FM, and VFW finalized the manuscript. VFW acquired funding and coordinated the study. All authors read and approved the final version of the manuscript.

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Conflict of Interest: The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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