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1	A rapid radiochemical filter paper assay for determination of Hexokinase activity and affinity for
2	glucose-6-phosphate (G6P)
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19 *Abstract*

20 Glucose phosphorylation by hexokinase (HK) is a rate-limiting step in glucose metabolism. Regulation 21 of HK includes feedback inhibition by its product glucose-6-phosphate (G6P) and mitochondria 22 binding. HK affinity for G6P is difficult to measure because its natural product (G6P) inhibits enzyme 23 activity. HK phosphorylates several hexoses and we have taken advantage of the fact that 2-24 deoxyglucose-6-phosphate (2DG-6P) does not inhibit HK activity. By this we have developed a new 25 method for rapid radiochemical analysis of HK activity with 2-deoxyglucose (2-DG) as substrate, 26 which allows control of the concentrations of G6P to investigate HK affinity for inhibition by G6P. We verified that 2DG serves as a substrate for the HK reaction with linear time and concentration 27 dependency as well as expected V_{max} and K_M . This is the first simple assay evaluating feedback 28 29 inhibition of HK by its product G6P and provides a unique technique for future research evaluating the regulation of glucose phosphorylation under various physiological conditions. 30

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32 *New & Noteworthy*

Traditionally, hexokinase activity has been analyzed spectrophotometrically where the product
formation of G6P is analyzed by an indirect reaction coupled to NADPH formation during conversion
of G6P to 6-P gluconolactone. By nature this assay prevents measurements of HK affinity for inhibition
by G6P. We have developed a rapid radiochemical filter paper assay to study HK affinity for G6P by
use of radiolabeled 2-DG as substrate, to study physiological regulation of HK affinity for G6Pinduced inhibition.

Introduction. Glucose uptake in various tissues is regulated by glucose delivery, glucose transport 40 41 across the membrane and intracellular phosphorylation (32). Although glucose phosphorylation can be 42 limiting for glucose uptake under various physiological conditions (2), the regulation of glucose 43 phosphorylation has received surprisingly little attention. Hexokinase (HK) catalyzes the irreversible 44 reaction converting glucose to glucose 6-phosphate (G6P). Glucose phosphorylation is a potential ratelimiting step for glucose metabolism and Mammalians express four different isoforms of hexokinase 45 46 (hexokinase I-IV) (37) while skeletal muscle expresses HK I and II (6, 11). HK is inhibited 47 allosterically by G6P with a K_i for G6P in the physiological range (37). It has been known for many years that HK binds to mitochondria and it was more recently reported that HK II is phosphorylated by 48 PKB at Thr⁴⁷³ (26). The translocation of HK to the mitochondria reduces the ability of G6P to inhibit 49 enzyme activity. However, the fact that the product inhibits HK activity makes is difficult to study the 50 physiological regulation of HK by G6P. 51

HK activity can easily be measured spectrophotometrically or fluorometrically (28). In this 52 53 method G6P-dehydrogenase (G6P-DH) in the assay buffer converts the formed G6P to 6-54 Phosphogluconolactone in a reaction coupled to conversion of NADP⁺ to NADPH; production of NADPH is measured to describe HK activity (28). Inclusion of G6P-DH in these methods is required 55 56 for the formation of NADPH, the presence of G6P-DH also prevents the accumulation of G6P which otherwise would exert a negative feedback on HK enzyme activity. Radiolabeled glucose or ATP can 57 58 also be used to measure HK activity where glucose and G6P can easily be separated. Sanderson et al. 59 used ¹⁴C-labelled glucose to measure HK activity and excluded G6P-DH from the assay buffer. In this study the authors demonstrated less inhibition of HK activity as G6P accumulated after insulin 60 stimulation in skeletal muscles from lean but not from obese Zucker rats (28). This suggests that insulin 61

decreases HK affinity for G6P-induced inhibition, but the method does not allow to calculate K_i
(inhibitory constant that is reflective of the binding affinity) for G6P-induced inhibition of HK. These
data suggest that HK affinity for G6P is regulated in skeletal muscles and highlights that methodologies
to investigate HK affinity for G6P are required.

Hexokinase phosphorylates a number of hexoses like 2-deoxyglucose (2-DG), but only G6P 66 and a few other hexose 6-phosphates inhibit HK activity (36). Importantly, 2-deoxyglucose-6-67 phosphate (2-DG-6P) does not inhibit HK activity (37). Thus, 2-DG serves as ideal substrate for HK 68 activity assay where the sensitivity for inhibition by G6P can be studied. Here we present a novel 69 radiochemical filter paper assay for rapid determination of HK activity and affinity for inhibition by 70 G6P. This method may not only be applicable for muscle physiology but also for cancer research as the 71 72 malignant phenotype of cancer cells is associated with increased hexokinase expression and glycolytic flux (20). 73

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75 Materials and methods

Experiments were conducted in Norway and Denmark. Experiments in Norway were approved by
Norwegian authorities and experimental procedures in Denmark by Danish and Animal Experimental
Inspectorate. All experiments were conducted in compliance with the European Union convention for
protection of vertebra animals used for scientific purposes (Council of Europe 123, Strasbourg, France,
1985).

81 Animals. In Norway male Wistar rats were purchased from B & K Universal (Nittedal, Norway). In

82 Denmark male Wistar rats and female C57B6 mice were purchased from Taconic (Ejby, Denmark).

Animals were kept in a room with 12:12-h light-dark cycle (light on/off, 6:00 AM/6:00 PM) and had
free access to water and chow.

85 *Tissue preparation and muscle incubation*. Animals were anesthetized by intraperitoneal injection of pentobarbital sodium (10 mg/100 g body wt) and tissues were dissected. Tissue samples were 86 homogenized directly (fresh tissue). In some studies, epitrochlearis and soleus were contracted or 87 treated with insulin in vitro prior to analysis of HK activity. Information about experimental conditions 88 is given in figure and table legends. For in vitro studies, muscles were dissected and suspended at 89 approximate resting length in an incubation system containing Krebs-Henseleit buffer (KHB) as 90 previously described (1). After 30 min preincubation, muscles were either rested for 30 min, stimulated 91 by supraphysiological insulin concentration (10 mU/ml) or forced to contract by electrical stimulation. 92 Muscle contraction was induced by electrical stimulation for 30 min consisting of 200-ms trains (100 93 Hz, 0.2 ms pulse duration, 10 V) delivered every 2 s, a protocol that previously has been shown to 94 result in a robust increase in glucose uptake (1). During the entire incubation period the buffer was 95 96 maintained at 30°C and oxygenated with 95% O₂ and 5% CO₂. After incubation, muscles were 97 harvested, blotted on filter paper and immediately homogenized.

98 *Homogenization*. Tissue was homogenized in ice cold buffer containing 250 mM sucrose, 20 mM

99 Sodium-pyrophosphate, 5 mM HEPES (pH 7.4), 5 mM magnesium chloride, 1 mM DTT, 5% Dextran

100 70, 20 mM β -glycerophosphate, 10 mM sodium fluoride, 2 mM PMSF, 1 mM EGTA (pH 8.0), 10

101 μ g/ml Aprotinin, 10 μ g/ml Leupeptin, 2 mM sodium orthovanadate and 3 mM benzamidine for 2 x 30

s at 30 Hz using a TissueLyzer II (Qiagen, Hilden, Germany).

103 Homogenates were analyzed for total protein concentration by use of the bicinchoninic acid method

104 (Thermo Scientific, Waltham, MA, USA).

A detailed list providing product no. and stock concentrations for reagents used in the homogenization
 buffer is given in Supplemental I (<u>dx.doi.org/10.17504/protocols.io.xcifiue</u>).

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108 *Filter paper HK activity assay.* Experiments were initially performed in Eppendorf tubes and the 109 reaction buffer with homogenate was spotted on DE81 anion exchanger filter paper (Whatman, UK, #3685 915) and washed as described below. To optimize and make the assay more efficient, the assay 110 111 was changed to 96-wells PCR plates (Thermo Fisher Scientific, AB0600). Homogenates were diluted to ~ 1-2 mg protein/ml in homogenization buffer. 15 μ l sample was added to a 96-well PCR plate 112 placed on an ice cold aluminum plate. As controls 15 µl homogenization buffer, cold assay buffer 113 114 (buffer blank), a standard pool and a blank sample (sample blank) was added to every assay plate. Kinase activity was started by adding 15 µl assay buffer in 1 sec intervals with a mutichannel pipette 115 while the aluminum plate was placed on a 30°C heating block. The assay buffer contained 50 mM TEA 116 (pH 7.4), 5 mM ATP, 5 mM magnesium chloride, 10 mM creatine phosphate, 2.4 µCi/ml ¹⁴C-2-117 deoxyglucose or ³H-2-deoxyglucose (Perkin Elmer), 2 mM 2DG, 0.05% BSA, 7 U/ml creatine kinase. 118 119 In case HK activity was evaluated in the presence of G6P or 2DG6P, various concentrations were 120 added to the assay buffer (see results for details). Kinase reaction was stopped at 30 °C after 15-25 min (depending on tissue and protein amount in the reaction) by adding 30 µl (1 vol) 90 % EtOH. The PCR 121 plate was kept under gentle agitation (manual agitation) at room temperature for 30 sec to ensure full 122 mixture and hence complete stop of kinase reaction. 20 μ l were spotted in duplicates by a multichannel 123 pipette on DE81 filter paper. The filter paper was subsequently washed 3 x 20 min in 300 ml 60% 124 125 EtOH in a plastic box (31cm x 22 cm x 6 cm) to remove non-phosphorylated 2DG and dried in a fume 126 hood. In order to determine specific activity, assay buffer $(2 \times 1 \mu l \text{ and } 2 \times 2 \mu l)$ was spotted on the

dried filter paper and added in vials containing 4 ml scintillation fluid. Finally the dried DE81 paper

128 was wrapped in plastic wrap and placed on a STORM cassette for 48 hours and analyzed on a STORM

129 Phosphoimager. Hexokinase activity was calculated as amount of ¹⁴C -2DG6 or ³H-2-DG6 captured in

the DE81 filter paper and given relative to specific activity of the assay buffer. A step-by-step

description of the assay is given in Supplemental II (<u>dx.doi.org/10.17504/protocols.io.xckfiuw</u>).

132 *Comparison of fluorometric and radiochemical assay for determination of HK activity.*

Tissue was homogenized as described above and analyzed for HK activity by the new radiochemicalfilter paper assay and compared to the well-established fluorometric method.

135 For fluorometric determination of HK activity, 20 µl sample was loaded as triplicate on a microtiter

136 plate and 200 µl reaction mixture was added. The reaction mixture contained 200 mM Tris buffer (Tris

HCl, Calbiochem 648313 and Tris Base, Calbiochem 648311) (pH 8.0), 10 mM magnesium chloride

138 (Sigma M2670), 12.5 mM Dithiothreitol (DTT) (Sigma D0362), 25 mM Glucose (Sigma G8270), 1

139 mM NADP, 11 U/ml G6PDH. The reaction was started by adding 10 µl ATP (100mM) (Sigma A

140 3377) and absorbance was measured at 340 nm as described by Lowry & Passoneau (18).

141 *Effect of exercise training on HK activity:* Quadriceps muscle samples from female WT mice

142 (SV129/C57B6 mixed background) from a previous study (9) were investigated for HK activity by use

143 of the new radiochemical filter paper assay and compared to the fluorometric method. Briefly, mice

144 were housed in single cages for 4 weeks without (sedentary) or with (training) access to a running

145 wheel. Mice were euthanized by cervical dislocation and muscle tissue was frozen in liquid nitrogen

and stored at -80 °C. We have previously investigated these samples by western blotting and reported a

147 ~3 fold increase in training-induced HKII protein expression (9).

148 *Results*.

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isolated by binding to the filter paper (Figure 1). Initially we compared the new radiochemical filter 150 paper assay to the established fluorometric method for determination of HK activity (Table 1). 151 Immediately after euthanization, rat brain, heart, soleus muscle and extensor digitorum longus (EDL) 152 muscle were homogenized. One portion of the homogenate was analyzed by the fluorometric assay 153 while another portion from the same homogenate sample was analyzed by the new radiochemical filter 154 155 paper assay. This enabled for direct comparison between these methods and revealed that the new filter paper assay yields comparable results to the established fluorometric assay (Table 1). 156 Based on multiple analyses intra-assay variation and inter-assay variation were calculated to be 157 5-10% and 10-16%, respectively. These were found to be comparable to other enzymatic assay such as 158 the GS-activity assay. Collectively these data demonstrate that the filter paper assay shows sensitivity 159 for detection of radiolabeled G6P that is comparable to the established fluorometric assay. 160 The fact that HK activity is inhibited by its product G6P precludes the use of glucose as 161 substrate for determination of HK affinity for G6P. However, hexokinase phosphorylates a number of 162 163 hexoses that do not inhibit HK activity and hence may serve as candidates to study alterations in affinity of HK activity for product inhibition by G6P (37). Therefore, we investigated 2-DG as a 164 substrate for the filter paper assay and compared it to glucose (Figure 2A). HK activity in rat heart was 165 166 assayed in the presence of increasing concentrations (0, 0.025, 0.05, 0.1, 0.2, 0.5, 1, 2 and 5 mM) of 2-DG and glucose, respectively (Figure 2A). Kinetic parameters were calculated by Lineweaver Burk 167 Plots (Figure 2B-C) and are given in Table 2. Hexokinase enzyme reaction velocity was substantially 168

In the new radiochemical filter paper assay glucose phosphorylated by HK to G6P will be

169	higher using 2-DG compared to glucose with a maximal velocity (V_{max}) of 14 and 6 μ mol/g
170	protein/min, respectively. K _M values derived from Lineweaver-Burk plot revealed a higher value for 2-
171	DG compared to glucose (0.67 vs. 0.16 mM). These findings do not preclude kinetic studies of HK
172	activity with 2-DG as a substrate; instead the markedly higher HK activity in light of a higher K_M
173	demonstrate that 2-DG serves as optimal substrate to study kinetic properties of HK. Furthermore,
174	product formation (2-DG6P) in rat heart increased linearly in a time- and concentration-dependent
175	manner (Figure 3A-B), validating the experimental settings of the radiochemical filter paper assay with
176	2-DG as a substrate.

Regular exercise training leads to a substantial increase in HK II expression and activity (35). In 177 order to verify that the new radiochemical filter paper assay is capable to detect alterations in HK 178 179 expression, we analyzed muscle sample from a previous mouse exercise training study (9). Determination of HK by the fluorometric and filter paper assay revealed a comparable increase in HK 180 activity by exercise training (~2.7 fold increase for fluorometric assay and ~2.5 fold increase for the 181 filter paper assay) (Figure 3C). These data are in accordance with the markedly higher levels of HK 182 183 expression we previously have reported (9) and demonstrate that the filter paper assay shows sensitivity for detecting differences in HK activity. 184

Given that majority of previous studies have investigated HK activation in frozen tissue, we tested whether the radiochemical filter paper assay also can be used to investigate HK activity in frozen tissue (Figure 3D). Rat soleus muscles were split in to two portions; one being homogenized as frozen tissue and the other part being homogenized as fresh tissue. Homogenate from frozen and fresh tissue was analyzed for HK activity by the fluorometric assay and the new filter paper assay. These data demonstrate that the radiochemical filter paper assay can be used to investigate HK activity in frozen tissue, albeit the HK activity generally tended to be a few percent lower in homogenate from frozen
tissue. Previous studies have reported HK binding to mitochondria as a potential mechanism for
reducing affinity for G6P-mediated inhibition (20, 26). In an attempt to maintain this interaction we
decided to perform subsequent analyses on fresh tissue.

Next, we studied HK activity in the presence of increasing concentrations (0, 0.1, 0.2, 0.5 and 1 mM) of 2-DG6P and G6P (Figure 4). This experiment reveals that increasing levels of G6P inhibit HK activity with a 50 % reduction (K_i) at ~0.2 mM G6P, while a similar increase in 2-DG6P in the assay buffer has no effect on HK activity. These findings verify that radiolabeled 2-DG can be used as substrate without that the formed 2-DG6P exerts a negative feedback on HK activity.

When assayed in the presence of increasing G6P concentrations, the physiological regulation of HK 200 201 affinity for G6P may be detected. Insulin and muscle contraction serve as main physiological stimuli to increase muscle glucose uptake and a subsequent rise in intracellular G6P levels (2, 14). Despite 202 elevated G6P levels skeletal glucose flux increases, suggesting that the negative feedback G6P exerts 203 204 on HK activity is antagonized under these physiological conditions. In order to investigate whether 205 sensitivity of HK for G6P underlies physiological regulation rat epitrochlearis muscles were stimulated ex vivo by insulin (10 mU/ml) or forced to contract for 30 min by electrical stimulation (Figure 5). 206 207 Maximal hexokinase activity (~1.8 µmol/g protein/min) was observed when assayed in the absence of 208 G6P with no effect of insulin stimulation or muscle contraction. HK activity decreased with increasing 209 G6P levels in a similar pattern in all three conditions and with a half-maximal inhibition (K_i) around 210 0.2 mM G6P. Thus, under the present experimental conditions HK affinity for G6P remained unaffected by insulin stimulation and muscle contraction. 211

213

214 Discussion

215 Hexokinase is the classical example of an enzyme that is inhibited by its product (G6P) in a feedback 216 dependent manner. However, regulation of HK activity is complex and HK translocates to 217 mitochondria and becomes phosphorylated (26). Translocation of HK to mitochondria has obtained large interest, but the effect of this translocation on HK activity is poorly understood. There are 218 219 indications that translocation of HK to the mitochondria reduces affinity for G6P-induced inhibition, but such analyses are hampered by the fact that HK is inhibited by its natural product G6P (37). Other 220 proteins are regulated by G6P; e.g. the major regulatory mechanism for glycogen synthase (GS) is by 221 222 changes in affinity for activation by G6P. This occurs via phosphorylation/dephosphorylation mechanisms (15, 17, 25). GS activation is normally reported as fractional activity, where activity at a 223 physiological concentration of G6P is related to total activity measured with a high concentration of 224 G6P sufficient to activate GS completely (12). Our idea was to establish a comparable assay where HK 225 activities are measured without G6P (total activity) and with a physiological concentration of G6P to 226 227 investigate changes in HK affinity for G6P-induced inhibition. Therefore, we have developed a simple 228 and rapid method for measurement of G6P-induced inhibition of HK by using 2-deoxyglucose (2-DG) as substrate to avoid inhibition by the product. This method can be used to evaluate regulation of HK 229 activity in various cell types and under various experimental conditions. 230

Here we present a rapid radiochemical filter paper assay that reveals HK activity measurements with a linear dose and time dependency comparable to the well-established fluorometric assay. Moreover, the use of radiolabeled 2-DG as a substrate in the filter paper assay allows to study the

sensitivity of HK for inhibition by its product G6P. We could demonstrate that increasing levels of G6P
but not 2-DG6P inhibit HK activity, providing a model to study the regulation of HK sensitivity for
G6P under various physiological settings.

Intracellular G6P serves as central regulator for skeletal muscle glycogen synthesis and should 237 be considered together with the degree of dephosphorylation of glycogen synthase when evaluating the 238 239 rate of glycogen synthesis (30). On the other hand G6P exerts a negative feedback on HK and the intracellular concentration of G6P must therefore be balanced between activation of glycogen synthase 240 and inhibition of HK (19). We observed a half maximal inhibition (K_i) of HK activity at ~0.2 mM G6P, 241 suggesting that HK was potently inhibited by levels of G6P thought to exist in skeletal muscle *in vivo*. 242 Thus, previous studies have reported G6P concentrations within the range of 0.2 - 0.3 mM under 243 resting conditions and increase above 1 mM in response to stimulation by insulin and muscle 244 contraction (14, 19, 23, 24). The G6P concentrations applied in the current HK assay (0.1-1 mM) 245 therefore reflect what could be expected in the physiological range. Increased glucose uptake and a 246 247 subsequent rise in intracellular G6P associated with insulin stimulation and muscle contraction should 248 consequently increase inhibition of hexokinase further, yet an increased flux is observed in vivo (2, 4, 27). This suggests the presence of mechanisms that uncouple the negative feedback associated G6P 249 250 levels and HK activity in vivo. In contrast to our hypothesis, HK sensitivity for G6P in response to 251 insulin stimulation and muscle contractions remained unaffected under the experimental conditions in 252 the present study. Sanderson et al (28) have previously reported that insulin reduced G6P-induced 253 inhibition in skeletal muscles, and other studies have proposed association of HK with mitochondrial membrane as a mechanism to regulate HK activity (3, 21, 37). Thus, once HK I and II are bound to 254 255 outer mitochondrial membrane inhibition of enzyme activity by G6P is antagonized. It has been

reported that insulin stimulation (3, 37) and muscle contraction (13, 33) increase binding of HK II to 256 257 the mitochondrial fraction. We used a homogenization buffer (containing 5% dextran 70 and 5 mM 258 $MgCl_2$) and a procedure (brief homogenization of fresh tissue) that previously has been shown to 259 maintain the integrity of the mitochondria (7, 8, 28, 34). Nevertheless we were unable to observe a 260 regulation of HK inhibition in response to insulin stimulation or muscle contraction. Insulin and 261 exercise have been shown to increase expression of HKII in human skeletal muscle whereas HKI 262 remains largely unaffected in response to these stimuli (16, 31). Notably this isoform specific 263 regulation was associated with increased HKII activity in soluble but not in the particulate fraction, highlighting the importance subcellular regulation when investigating the regulation of HK activity. In 264 the present study HK enzyme activity was not investigated isoform specific in cellular sub-fractions, so 265 266 we cannot rule out the possibility that regulation of HKII activity may have been overseen due to the crude measurement in the present study. 267

Hexokinase binds to the mitochondria (VDAC) via it N-terminal end and mutating the fifth 268 269 amino acid histidine to proline completely abolish HK-2 binding to the mitochondria (5). The 270 interaction between HK2 and VDAC is not characterized, but it tempting to speculate that the binding is rather loose and may be disrupted during homogenization. However, mitochondria can be isolated 271 272 with attached HK (22) and it could therefore be speculated that isolation of mitochondrial fraction is 273 required to study these alterations in G6P-mediated inhibition of HK activity. Indeed, Van Houten and 274 colleagues reported an increase in HK activity in the mitochondrial fraction but not total fraction when 275 rat gastrocnemius muscles were analyzed following swimming exercise until exhaustion (13). We suggest that other homogenization procedures or studies in saponised muscle bundles should be 276 investigated. 277

The interaction between HKII and mitochondria (VDAC) has large interest in cancer research 278 279 and emerged as a drug target for cancer therapies (29) and disruption of HKII-mitochondria interaction with peptides corresponding to 15 aa of the N-terminal of HKII disrupt the interaction and triggers 280 281 apoptosis (38). The mechanism by which this disruption triggers apoptosis remain unknown and we 282 believe assaying HK affinity for G6P-induced inhibition will be most useful to solve this question. 283 HKII binding to mitochondria has also been suggested as key aspect in cardiac protection (10). It has been demonstrated in cardiomyocytes that HK is phosphorylated on Thr⁴⁷³ by PKB, a regulatory 284 285 mechanisms that was found to be associated with translocation of HK to the mitochondria and was associated with reduced affinity for G6P-mediated inhibition (26). This adds an additional regulatory 286 287 mechanism for control of HK inhibition by G6P, although the presence in intact mature tissue such as 288 skeletal muscle remains to be demonstrated. A simple method to measure HK affinity for G6P-induced inhibition will be a useful tool to address such question. 289

This is the first assay that takes advantage of 2-DG as substrate to study the sensitivity of HK activity for G6P. Abnormal regulation of HK activity has previously been reported in disease state such as muscle insulin resistance in type 2 diabetes (28) or the increased glycolytic flux present in malignant cancer cells (20). The presented method provides a valuable tool to study the regulation of HK inhibition by G6P under these conditions. Future research could therefore apply this assay in order to clarify the physiological regulation of HK affinity in diseased states such as T2D and cancer.

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403 Figure legends

Figure 1: Assay reaction by the new established radiochemical filter paper assay that was 404 405 developed to study hexokinase sensitivity for inhibition by G6P. A rapid radiochemical filter paper assay for measurement of hexokinase (HK) activity and affinity for 406 G6P was developed. This assay allows for the simple separation of phosphorylated form of the 407 radiolabeled tracer ($[^{14}C]$ - Glucose-phosphate or $[^{3}H]$ -2-Deoxyglucose-phosphate) from the 408 unphosphorylated form ($[^{14}C]$ - Glucose and $[^{3}H]$ -2-deoxyglucose). 409 In this assay conversion of radiolabeled 2-Deoxyglucose (2DG) to 2-Deoxyglucose-6-phosphate (2-410 DG-6-phosphate) is catalyzed by HK. 2DG-6-phosphate binds to DE81 ion-exchange filter paper 411 (Whatman, Maidstone, UK) and non-phosphorylated 2-DG can be separated by washing the filter paper 412 in excess 60% EtOH. Importantly, 2-DG-6-phosphate does not inhibit HK activity and 2-DG is 413 therefore a perfect substrate for studying the HK sensitivity for G6P. Thus, HK affinity can easily be 414 investigated by performing the radiochemical filter paper assay in the presence of varying G6P 415 416 concentration.

417

418 Figure 2: Comparison of substrates for radiochemical filter paper assay to study hexokinase

419 **activity.** Hexokinase activity in 6 homogenates derived from a single rat heart was measured in the

420 presence of increasing concentration (0, 0.025, 0.05, 0.1, 0.2, 0.5, 1, 2 and 5 mM) of glucose for 14 C-

- 421 glucose and 2-deoxyglucose (2-DG) for ³H-2-deoxyglucose, respectively (A). Data are representing
- 422 means and variation (SEM) for 6 homogenates obtained from the same heart tissue. Lineweaver-Burk
- 423 Plots were used to calculate V_{max} and K_M for ³H-2-DG and ¹⁴C-Glucose (B-C) and are given in Table 2.

424

425 Figure 3. Sensitivity of the radiochemical filter paper assay

Hexokinase activity in a single rat heart homogenate was measured over time (A) and in the presence
of increasing sample/protein concentration for 2 min (B) by use of the radiochemical filter paper assay
with 2-DG as substrate and ³H-2DG as tracer. Product formation of 2-DG-6P was calculated. Each data
point represents a triplicate.

The effect of exercise training on HK activity was investigated in mouse muscle from a previous study 430 (9). Quadriceps muscle from sedentary mice and 4-week trained mice (access to running wheel) were 431 compared by the fluorometric and the radiochemical filter paper assay (C). Data from 4 sedentary and 4 432 trained mice are presented as means \pm SEM (n=4). For direct comparison between fresh and frozen 433 tissue, rat soleus muscles were split into two pieces and homogenized as fresh and as frozen tissue 434 respectively (D). HK activity was subsequently determined by fluorometric method and the 435 radiochemical filter paper assay (C). Data represent means \pm SEM for 4 soleus muscles that were split 436 437 into two pieces and investigated by by fluorometric method and the radiochemical filter paper assay 438 (n=4). T-test within fluorometric assay and filter paper assay was used to investigate for the effect of exercise training and fresh/frozen muscle respectively. *** p<0.001 for effect of exercise training. 439

440 Figure 4. Hexokinase activity in the presence of increasing 2DG-6P and G6P concentrations.

Hexokinase activity in mouse heart was analyzed by the radiochemical filter paper assay with ¹⁴C-2DG as a substrate and assayed in the presence of increasing 2-DG6P or G6P concentrations (0, 0.1, 0.2, 0.5, 1 mM), respectively. Data represent heart samples from 8 mice and are given as means \pm SEM (n=8).

445 Figure 5. Physiological regulation hexokinase sensitivity for G6P-mediated inhibition. Rat

446 epitrochlearis muscles were isolated and suspended in an *ex vivo* incubation system. Muscles were

rested (basal non-stimulated control), stimulated by insulin (10 mU/ml) or forced to contract for 30min.

448 Immediately after incubation, muscles were homogenized and analyzed for HK activity. Sensitivity of

HK activity for G6P was analyzed with use radiochemical filter paper assay with 14 C-2DG as a

450 substrate assayed in the presence of increasing G6P concentrations (0, 0.1, 0.2, 0.5, 1 mM).

451 Data represent values from 8 rested control muscle, 8 insulin-stimulated muscles and 8 electrically

452 contracted muscles and are given as means+/-SEM (n=8).

453 Table 1: Comparison of fluorometric and radiochemical assay for hexokinase activity.

454 Rat brain, heart, soleus muscle and EDL muscle were homogenized immediately after cervical

dislocation. HK activity in homogenate samples was determined by the established fluorometric assay

456 (formation of NADH) and compared to the newly established radiochemical filter paper assay (^{14}C -

457 2DG 6-phosphate). Activity is given as µmol/g protein/min.

458 Data present HK activity in different tissues from 8 Wistar rats (n=8) and are presented as means and 459 the variation among samples is given as SEM (n=8). T-tests revealed no statistical differences between

460 fluorometric and radiochemical assay for measurement of HK activity.

461 Table 2: Kinetic parameters for 2-deoxyglucose (2DG) and Glucose

462 Lineweaver-Burk Plots were used to calculate V_{max} and K_M for ³H-2-DG and ¹⁴C-Glucose (Figure 2B-463 C) and are given in Table 2.

Figure 1

Assay reaction

¹⁴C-2-DG + ATP $\xrightarrow{\text{HK}}$ ¹⁴C-2DG 6-phosphate + ADP + different concentrations of G6P

Table 1

	Flurometric assay	Filter paper assay
Brain	28.76 ± 2.87	26.01 ± 1.31
Heart	10.32 ± 1.04	11.63 ± 1.68
Soleus	2.31 ± 0.05	2.21 ± 0.17
EDL	1.23 ± 0.14	1.14 ± 0.29

Figure 2



В ³H-2DG: Lineweaver-Burk plot 0,4 y = 0,0478x + 0,07090,3 1/V_{0,2} 0,1 0 -1 -0,1 3 5 -5 1 1/[S]

-0,2

С

¹⁴C-Glucose: Lineweaver-Burk plot



Table 2: Kinetic parameters for 2DG and Glucose

	V _{max}	К _м
	(µmol/g protein/min)	(mM)
³ H-2DG	14	0.67
¹⁴ C-Glucose	6	0.16

Figure 3:

Α



В

Product formation - Effect of protein concentration



С



D



Figure 4



Figure 5

